METHODS FOR QUANTITATING SMALL RNA MOLECULES

FIELD OF THE INVENTION

The present invention relates to methods of amplifying and quantitating small RNA molecules.

BACKGROUND OF THE INVENTION

RNA interference (RNAi) is an evolutionarily conserved process that functions to inhibit gene expression (Bernstein et al. (2001), *Nature 409*:363-6; Dykxhoorn et al. (2003) *Nat. Rev. Mol. Cell. Biol. 4*:457-67). The phenomenon of RNAi was first described in *Caenorhabditis elegans*, where injection of double-stranded RNA (dsRNA) led to efficient sequence-specific gene silencing of the mRNA that was complementary to the dsRNA (Fire et al. (1998) *Nature 391*:806-11). RNAi has also been described in plants as a phenomenon called post-transcriptional gene silencing (PTGS), which is likely used as a viral defense mechanism (Jorgensen (1990) *Trends Biotechnol. 8*:340-4; Brigneti et al. (1998) *EMBO J. 17*:6739-46; Hamilton & Baulcombe (1999) *Science* 286:950-2).

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An early indication that the molecules that regulate PTGS were short RNAs processed from longer dsRNA was the identification of short 21 to 22 nucleotide dsRNA derived from the longer dsRNA in plants (Hamilton & Baulcombe (1999) Science 286:950-2). This observation was repeated in Drosophila embryo extracts where long dsRNA was found processed into 21-25 nucleotide short RNA by the RNase III type enzyme, Dicer (Elbashir et al. (2001) Nature 411:494-8; Elbashir et al. (2001) EMBO J. 20:6877-88; Elbashir et al. (2001) Genes Dev. 15:188-200). These observations led Elbashir et al. to test if synthetic 21-25 nucleotide synthetic dsRNAs function to specifically inhibit gene expression in Drosophila embryo lysates and mammalian cell

culture (Elbashir et al. (2001) *Nature 411*:494-8; Elbashir et al. (2001) *EMBO J.* 20:6877-88; Elbashir et al. (2001) *Genes Dev. 15*:188-200). They demonstrated that small interfering RNAs (siRNAs) had the ability to specifically inhibit gene expression in mammalian cell culture without induction of the interferon response.

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These observations led to the development of techniques for the reduction, or elimination, of expression of specific genes in mammalian cell culture, such as plasmid-based systems that generate hairpin siRNAs (Brummelkamp et al. (2002) *Science 296*:550-3; Paddison et al. (2002) *Genes Dev. 16*:948-58; Paddison et al. (2002) *Proc. Natl. Acad. Sci. U.S.A. 99*:1443-8; Paul et al. 2002) *Nat. Biotechnol. 20*:404-8). siRNA molecules can also be introduced into cells, *in vivo*, to inhibit the expression of specific proteins (see, e.g., Soutschek, J., et al., *Nature 432* (7014):173-178 (2004)).

siRNA molecules have promise both as therapeutic agents for inhibiting the expression of specific proteins, and as targets for drugs that affect the activity of siRNA molecules that function to regulate the expression of proteins involved in a disease state. A first step in developing such therapeutic agents is to measure the amounts of specific siRNA molecules in different cell types within an organism, and thereby construct an "atlas" of siRNA expression within the body. Additionally, it will be useful to measure changes in the amount of specific siRNA molecules in specific cell types in response to a defined stimulus, or in a disease state.

Short RNA molecules are difficult to quantitate. For example, with respect to the use of PCR to amplify and measure the small RNA molecules, most PCR primers are longer than the small RNA molecules, and so it is difficult to design a primer that has significant overlap with a small RNA molecule, and that selectively hybridizes to the small RNA molecule at the temperatures used for primer extension and PCR amplification reactions.

SUMMARY OF THE INVENTION

In one aspect, the present invention provides methods for amplifying a microRNA molecule to produce cDNA molecules. The methods include the steps of: (a) producing a first DNA molecule that is complementary to a target microRNA molecule using primer extension; and (b) amplifying the first DNA molecule to produce amplified DNA molecules using a universal forward primer and a reverse primer. In some embodiments of the method, at least one of the forward primer and the reverse primer comprise at least one locked nucleic acid molecule. It will be understood that, in the practice of the present

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invention, typically numerous (e.g., millions) of individual microRNA molecules are amplified in a sample (e.g., a solution of RNA molecules isolated from living cells).

In another aspect, the present invention provides methods for measuring the amount of a target microRNA in a a sample from a living organism. The methods of this aspect of the invention include the step of measuring the amount of a target microRNA molecule in a multiplicity of different cell types within a living organism, wherein the amount of the target microRNA molecule is measured by a method including the steps of: (1) producing a first DNA molecule complementary to the target microRNA molecule in the sample using primer extension; (2) amplifying the first DNA molecule to produce amplified DNA molecules using a universal forward primer and a reverse primer; and (3) measuring the amount of the amplified DNA molecules. In some embodiments of the method, at least one of the forward primer and the reverse primer comprise at least one locked nucleic acid molecule.

In another aspect, the invention provides nucleic acid primer molecules consisting of sequence SEQ ID NO:1 to SEQ ID NO: 499, as shown in TABLE 1, TABLE 2, TABLE 6 and TABLE 7. The primer molecules of the invention can be used as primers for detecting mammalian microRNA target molecules, using the methods of the invention described herein.

In another aspect, the present invention provides kits for detecting at least one mammalian target microRNA, the kits comprising one or more primer sets specific for the detection of a target microRNA, each primer set comprising (1) an extension primer for producing a cDNA molecule complementary to a target microRNA, (2) a universal forward PCR primer for amplifying the cDNA molecule and (3) a reverse PCR primer for amplifying the cDNA molecule. The extension primer comprises a first portion that hybridizes to the target microRNA molecule and a second portion that includes a hybridization sequence for a universal forward PCR primer. The reverse PCR primer comprises a sequence selected to hybridize to a portion of the cDNA molecule. In some embodiments of the kit, at least one of the universal forward and reverse primers include at least one locked nucleic acid molecule. The kits of the invention may be used to practice various embodiments of the methods of the invention.

The present invention is useful, for example, for quantitating specific microRNA molecules within different types of cells in a living organism, or, for example, for

measuring changes in the amount of specific microRNAs in living cells in response to a stimulus (e.g., in response to administration of a drug).

BRIEF DESCRIPTION OF THE DRAWINGS

The foregoing aspects and many of the attendant advantages of this invention will become more readily appreciated as the same become better understood by reference to the following detailed description, when taken in conjunction with the accompanying drawings, wherein:

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FIGURE 1 shows a flow chart of a representative method of the present invention; FIGURE 2 graphically illustrates the standard curves for assays specific for the detection of microRNA targets miR-95 and miR-424 as described in EXAMPLE 3;

FIGURE 3A is a histogram plot showing the expression profile of miR-1 across a panel of total RNA isolated from twelve tissues as described in EXAMPLE 5;

FIGURE 3B is a histogram plot showing the expression profile of miR-124 across a panel of total RNA isolated from twelve tissues as described in EXAMPLE 5; and

FIGURE 3C is a histogram plot showing the expression profile of miR-150 across a panel of total RNA isolated from twelve tissues as described in EXAMPLE 5.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

In accordance with the foregoing, in one aspect, the present invention provides methods for amplifying a microRNA molecule to produce cDNA molecules. The methods include the steps of: (a) using primer extension to make a DNA molecule that is complementary to a target microRNA molecule; and (b) using a universal forward primer and a reverse primer to amplify the DNA molecule to produce amplified DNA molecules. In some embodiments of the method, at least one of the universal forward primer and the reverse primer comprises at least one locked nucleic acid molecule.

As used herein, the term "locked nucleic acid molecule" (abbreviated as LNA molecule) refers to a nucleic acid molecule that includes a 2'-O,4'-C-methylene-β-D-ribofuranosyl moiety. Exemplary 2'-O,4'-C-methylene-β-D-ribofuranosyl moieties, and exemplary LNAs including such moieties, are described, for example, in Petersen, M. and Wengel, J., *Trends in Biotechnology 21*(2):74-81 (2003) which publication is incorporated herein by reference in its entirety.

As used herein, the term "microRNA" refers to an RNA molecule that has a length in the range of from 21 nucleotides to 25 nucleotides. Some microRNA molecules (e.g., siRNA molecules) function in living cells to regulate gene expression.

Representative method of the invention. FIGURE 1 shows a flowchart of a representative method of the present invention. In the method represented in FIGURE 1, a microRNA is the template for synthesis of a complementary first DNA molecule. The synthesis of the first DNA molecule is primed by an extension primer, and so the first DNA molecule includes the extension primer and newly synthesized DNA (represented by a dotted line in FIGURE 1). The synthesis of DNA is catalyzed by reverse transcriptase.

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The extension primer includes a first portion (abbreviated as FP in FIGURE 1) and a second portion (abbreviated as SP in FIGURE 1). The first portion hybridizes to the microRNA target template, and the second portion includes a nucleic acid sequence that hybridizes with a universal forward primer, as described *infra*.

A quantitative polymerase chain reaction is used to make a second DNA molecule that is complementary to the first DNA molecule. The synthesis of the second DNA molecule is primed by the reverse primer that has a sequence that is selected to specifically hybridize to a portion of the target first DNA molecule. Thus, the reverse primer does not hybridize to nucleic acid molecules other than the first DNA molecule. The reverse primer may optionally include at least one LNA molecule located within the portion of the reverse primer that does not overlap with the extension primer. In FIGURE 1, the LNA molecules are represented by shaded ovals.

A universal forward primer hybridizes to the 3' end of the second DNA molecule and primes synthesis of a third DNA molecule. It will be understood that, although a single microRNA molecule, single first DNA molecule, single second DNA molecule, single third DNA molecule and single extension, forward and reverse primers are shown in FIGURE 1, typically the practice of the present invention uses reaction mixtures that include numerous copies (e.g., millions of copies) of each of the foregoing nucleic acid molecules.

The steps of the methods of the present invention are now considered in more detail.

<u>Preparation of microRNA molecules useful as templates</u>. microRNA molecules useful as templates in the methods of the invention can be isolated from any organism (e.g., eukaryote, such as a mammal) or part thereof, including organs, tissues, and/or individual cells (including cultured cells). Any suitable RNA preparation that includes microRNAs can be used, such as total cellular RNA.

RNA may be isolated from cells by procedures that involve lysis of the cells and denaturation of the proteins contained therein. Cells of interest include wild-type cells, drug-exposed wild-type cells, modified cells, and drug-exposed modified cells.

Additional steps may be employed to remove some or all of the DNA. Cell lysis may be accomplished with a nonionic detergent, followed by microcentrifugation to remove the nuclei and hence the bulk of the cellular DNA. In one embodiment, RNA is extracted from cells of the various types of interest using guanidinium thiocyanate lysis followed by CsCl centrifugation to separate the RNA from DNA (see, Chirgwin et al., 1979, Biochemistry 18:5294-5299). Separation of RNA from DNA can also be accomplished by organic extraction, for example, with hot phenol phenol/chloroform/isoamyl alcohol.

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If desired, RNase inhibitors may be added to the lysis buffer. Likewise, for certain cell types, it may be desirable to add a protein denaturation/digestion step to the protocol.

The sample of RNA can comprise a multiplicity of different microRNA molecules, each different microRNA molecule having a different nucleotide sequence. In a specific embodiment, the microRNA molecules in the RNA sample comprise at least 100 different nucleotide sequences. In other embodiments, the microRNA molecules of the RNA sample comprise at least 500, 1,000, 5,000, 10,000, 20,000, 30,000, 40,000, 50,000, 60,000, 70,000, 80,000 90,000, or 100,000 different nucleotide sequences.

The methods of the invention may be used to detect the presence of any microRNA. For example, the methods of the invention can be used to detect one or more of the microRNA targets described in a database such as "the miRBase sequence database" as described in Griffith-Jones et al. (2004), *Nucleic Acids Research 32*:D109-D111, and Griffith-Jones et al. (2006), *Nucleic Acids Research* 34: D140-D144, which is publically accessible on the World Wide Web at the Wellcome Trust Sanger Institute website at http://microrna.sanger.ac.uk/sequences/. A list of exemplary microRNA targets is also described in the following references: Lagos-Quintana et al., *Curr. Biol.* 12(9):735-9 (2002).

Synthesis of DNA molecules using microRNA molecules as templates. In the practice of the methods of the invention, first DNA molecules are synthesized that are complementary to the microRNA target molecules, and that are composed of an extension primer and newly synthesized DNA (wherein the extension primer primes the

synthesis of the newly synthesized DNA). Individual first DNA molecules can be complementary to a whole microRNA target molecule, or to a portion thereof; although typically an individual first DNA molecule is complementary to a whole microRNA target molecule. Thus, in the practice of the methods of the invention, a population of first DNA molecules is synthesized that includes individual DNA molecules that are each complementary to all, or to a portion, of a target microRNA molecule.

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The synthesis of the first DNA molecules is catalyzed by reverse transcriptase. Any reverse transcriptase molecule can be used to synthesize the first DNA molecules, such as those derived from Moloney murine leukemia virus (MMLV-RT), avian myeloblastosis virus (AMV-RT), bovine leukemia virus (BLV-RT), Rous sarcoma virus (RSV) and human immunodeficiency virus (HIV-RT). A reverse transcriptase lacking RNaseH activity (e.g., Superscript IIITM sold by Invitrogen, 1600 Faraday Avenue, PO Box 6482, Carlsbad, California 92008) is preferred in order to minimize the amount of double-stranded cDNA synthesized at this stage. The reverse transcriptase molecule should also preferably be thermostable so that the DNA synthesis reaction can be conducted at as high a temperature as possible, while still permitting hybridization of primer to the microRNA target molecules.

Priming the synthesis of the first DNA molecules. The synthesis of the first DNA molecules is primed using an extension primer. Typically, the length of the extension primer is in the range of from 10 nucleotides to 100 nucleotides, such as 20 to 35 nucleotides. The nucleic acid sequence of the extension primer is incorporated into the sequence of each, synthesized, DNA molecule. The extension primer includes a first portion that hybridizes to a portion of the microRNA molecule. Typically the first portion of the extension primer includes the 3'-end of the extension primer. The first portion of the extension primer typically has a length in the range of from 6 nucleotides to 20 nucleotides, such as from 10 nucleotides to 12 nucleotides. In some embodiments, the first portion of the extension primer has a length in the range of from 3 nucleotides to 25 nucleotides.

The extension primer also includes a second portion that typically has a length of from 18 to 25 nucleotides. For example, the second portion of the extension primer can be 20 nucleotides long. The second portion of the extension primer is located 5' to the first portion of the extension primer. The second portion of the extension primer includes at least a portion of the hybridization site for the universal forward primer. For example,

the second portion of the extension primer can include all of the hybridization site for the universal forward primer, or, for example, can include as little as a single nucleotide of the hybridization site for the universal forward primer (the remaining portion of the hybridization site for the forward primer can, for example, be located in the first portion of the extension primer). An exemplary nucleic acid sequence of a second portion of an extension primer is 5' CATGATCAGCTGGGCCAAGA 3' (SEQ ID NO:1).

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Amplification of the DNA molecules. In the practice of the methods of the invention, the fist DNA molecules are enzymatically amplified using the polymerase chain reaction. A universal forward primer and a reverse primer are used to prime the polymerase chain reaction. The reverse primer includes a nucleic acid sequence that is selected to specifically hybridize to a portion of a first DNA molecule.

The reverse primer typically has a length in the range of from 10 nucleotides to 100 nucleotides. In some embodiments, the reverse primer has a length in the range of from 12 nucleotides to 20 nucleotides. The nucleotide sequence of the reverse primer is selected to hybridize to a specific target nucleotide sequence under defined hybridization conditions. The reverse primer and extension primer are both present in the PCR reaction mixture, and so the reverse primer should be sufficiently long so that the melting temperature (Tm) is at least 50°C, but should not be so long that there is extensive overlap with the extension primer which may cause the formation of "primer dimers." "Primer dimers" are formed when the reverse primer hybridizes to the extension primer, and uses the extension primer as a substrate for DNA synthesis, and the extension primer hybridizes to the reverse primer, and uses the reverse primer as a substrate for DNA synthesis. To avoid the formation of "primer dimers," typically the reverse primer and the extension primer are designed so that they do not overlap with each other by more than 6 nucleotides. If it is not possible to make a reverse primer having a Tm of at least 50°C, and wherein the reverse primer and the extension primer do not overlap by more than 6 nucleotides, then it is preferable to lengthen the reverse primer (since Tm usually increases with increasing oligonucleotide length) and decrease the length of the extension primer.

The reverse primer primes the synthesis of a second DNA molecule that is complementary to the first DNA molecule. The universal forward primer hybridizes to the portion of the second DNA molecule that is complementary to the second portion of the extension primer which is incorporated into all of the first DNA molecules. The

universal forward primer primes the synthesis of third DNA molecules. The universal forward primer typically has a length in the range of from 16 nucleotides to 100 nucleotides. In some embodiments, the universal forward primer has a length in the range of from 16 nucleotides to 30 nucleotides. The universal forward primer may include at least one locked nucleic acid molecule. In some embodiments, the universal forward primer includes from 1 to 25 locked nucleic acid molecules. The nucleic acid sequence of an exemplary universal forward primer is set forth in SEQ ID NO:13.

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In general, the greater the number of amplification cycles during the polymerase chain reaction, the greater the amount of amplified DNA that is obtained. On the other hand, too many amplification cycles (e.g., more than 35 amplification cycles) may result in spurious and unintended amplification of non-target double-stranded DNA. Thus, in some embodiments, a desirable number of amplification cycles is between one and 45 amplification cycles, such as from one to 25 amplification cycles, or such as from five to 15 amplification cycles, or such as ten amplification cycles.

Use of LNA molecules and selection of primer hybridization conditions: hybridization conditions are selected that promote the specific hybridization of a primer molecule to the complementary sequence on a substrate molecule. With respect to the hybridization of a 12 nucleotide first portion of an extension primer to a microRNA, it has been found that specific hybridization occurs at a temperature of 50°C. Similarly, it has been found that hybridization of a 20 nucleotide universal forward primer to a complementary DNA molecule, and hybridization of a reverse primer (having a length in the range of from 12-20 nucleotides, such as from 14-16 nucleotides) to a complementary DNA molecule occurs at a temperature of 50°C. By way of example, it is often desirable to design extension, reverse and universal forward primers that each have a hybridization temperature in the range of from 50°C to 60°C.

In some embodiments, LNA molecules can be incorporated into at least one of the extension primer, reverse primer, and universal forward primer to raise the Tm of one, or more, of the foregoing primers to at least 50°C. Incorporation of an LNA molecule into the portion of the reverse primer that hybridizes to the target first DNA molecule, but not to the extension primer, may be useful because this portion of the reverse primer is typically no more than 10 nucleotides in length. For example, the portion of the reverse primer that hybridizes to the target first DNA molecule, but not to the extension primer, may include at least one locked nucleic acid molecule (e.g., from 1 to 25 locked nucleic

acid molecules). In some embodiments, two or three locked nucleic acid molecules are included within the first 8 nucleotides from the 5' end of the reverse primer.

The number of LNA residues that must be incorporated into a specific primer to raise the Tm to a desired temperature mainly depends on the length of the primer and the nucleotide composition of the primer. A tool for determining the effect on Tm of one or more LNAs in a primer is available on the Internet Web site of Exiqon, Bygstubben 9, DK-2950 Vedbaek, Denmark.

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Although one or more LNAs can be included in any of the primers used in the practice of the present invention, it has been found that the efficiency of synthesis of cDNA is low if an LNA is incorporated into the extension primer. While not wishing to be bound by theory, LNAs may inhibit the activity of reverse transcriptase.

Detecting and measuring the amount of the amplified DNA molecules: amplified DNA molecules can be detected and quantitated by the presence of detectable marker molecules, such as fluorescent molecules. For example, the amplified DNA molecules can be detected and quantitated by the presence of a dye (e.g., SYBR green) that preferentially or exclusively binds to double stranded DNA during the PCR amplification step of the methods of the present invention. For example, Molecular Probes, Inc. (29851 Willow Creek Road, Eugene, OR 97402) sells quantitative PCR reaction mixtures that include SYBR green dye. By way of further example, another dye (referred to as "BEBO") that can be used to label double stranded DNA produced during real-time PCR is described by Bengtsson, M., et al., Nucleic Acids Research 31(8):e45 (April 15, 2003), which publication is incorporated herein by reference. Again by way of example, a forward and/or reverse primer that includes a fluorophore and quencher can be used to prime the PCR amplification step of the methods of the present invention. The physical separation of the fluorophore and quencher that occurs after extension of the labeled primer during PCR permits the fluorophore to fluoresce, and the fluorescence can be used to measure the amount of the PCR amplification products. Examples of commercially available primers that include a fluorophore and quencher include Scorpion primers and Uniprimers, which are both sold by Molecular Probes, Inc.

Representative uses of the present invention: The present invention is useful for producing cDNA molecules from microRNA target molecules. The amount of the DNA molecules can be measured which provides a measurement of the amount of target microRNA molecules in the starting material. For example, the methods of the present

invention can be used to measure the amount of specific microRNA molecules (e.g., specific siRNA molecules) in living cells. Again by way of example, the present invention can be used to measure the amount of specific microRNA molecules (e.g., specific siRNA molecules) in different cell types in a living body, thereby producing an "atlas" of the distribution of specific microRNA molecules within the body. Again by way of example, the present invention can be used to measure changes in the amount of specific microRNA molecules (e.g., specific siRNA molecules) in response to a stimulus, such as in response to treatment of a population of living cells with a drug.

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Thus, in another aspect, the present invention provides methods for measuring the amount of a target microRNA in a multiplicity of different cell types within a living organism (e.g., to make a microRNA "atlas" of the organism). The methods of this aspect of the invention each include the step of measuring the amount of a target microRNA molecule in a multiplicity of different cell types within a living organism, wherein the amount of the target microRNA molecule is measured by a method comprising the steps of: (1) using primer extension to make a DNA molecule complementary to the target microRNA molecule isolated from a cell type of a living organism; (2) using a universal forward primer and a reverse primer to amplify the DNA molecule to produce amplified DNA molecules, and (3) measuring the amount of the amplified DNA molecules. In some embodiments of the methods, at least one of the forward primer and the reverse primer comprises at least one locked nucleic acid molecule. The measured amounts of amplified DNA molecules can, for example, be stored in an interrogatable database in electronic form, such as on a computer-readable medium (e.g., a floppy disc).

In another aspect, the invention provides nucleic acid primer molecules consisting of sequence SEQ ID NO:1 to SEQ ID NO: 499, as shown in TABLE 1, TABLE 2, TABLE 6 and TABLE 7. The primer molecules of the invention can be used as primers for detecting mammalian microRNA target molecules, using the methods of the invention described herein.

In another aspect, the present invention provides kits for detecting at least one mammalian target microRNA, the kits comprising one or more primer sets specific for the detection of a target microRNA, each primer set comprising (1) an extension primer for producing a cDNA molecule complementary to a target microRNA, (2) a universal forward PCR primer and (3) a reverse PCR primer for amplifying the cDNA molecule. The extension primer comprises a first portion that hybridizes to the target microRNA

molecule and a second portion that includes a hybridization sequence for a universal forward PCR primer. The reverse PCR primer comprises a sequence selected to hybridize to a portion of the cDNA molecule. In some embodiments of the kits, at least one of the universal forward and reverse primers includes at least one locked nucleic acid molecule.

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The extension primer, universal forward and reverse primers for inclusion in the kit may be designed to detect any mammalian target microRNA in accordance with the methods described herein. Nonlimiting examples of human target microRNA target molecules and exemplary target-specific extension primers and reverse primers are listed below in TABLE 1, TABLE 2 and TABLE 6. Nonlimiting examples of murine target microRNA target molecules and exemplary target-specific extension primers and reverse primers are listed below in TABLE 7. A nonlimiting example of a universal forward primer is set forth as SEQ ID NO: 13.

In certain embodiments, the kit includes a set of primers comprising an extension primer, reverse and universal forward primers for a selected target microRNA molecule that each have a hybridization temperature in the range of from 50°C to 60°C.

In certain embodiments, the kit includes a plurality of primer sets that may be used to detect a plurality of mammalian microRNA targets, such as two microRNA targets up to several hundred microRNA targets.

In certain embodiments, the kit comprises one or more primer sets capable of detecting at least one or more of the following human microRNA target templates: of miR-1, miR-7, miR-9*, miR-10a, miR-10b, miR-15a, miR-15b, miR-16, miR-17-3p, miR-17-5p, miR-18, miR-19a, miR-19b, miR-20, miR-21, miR-22, miR-23a, miR-23b, miR-24, miR-25, miR-26a, miR-26b, miR-27a, miR-28, miR-29a, miR-29b, miR-29c, miR-30a-5p, miR-30b, miR-30c, miR-30d, miR-30e-5p, miR-30e-3p, miR-31, miR-32, miR-33, miR-34a, miR-34b, miR-34c, miR-92, miR-93, miR-95, miR-96, miR-98, miR-99a, miR-99b, miR-100, miR-101, miR-103, miR-105, miR-106a, miR-107, miR-122, miR-122a, miR-124, miR-124a, miR-125a, miR-125b, miR-126, miR-126*, miR-127, miR-128a, miR-128b, miR-129, miR-130a, miR-130b, miR-132, miR-133a, miR-133b, miR-134, miR-135a, miR-135b, miR-136, miR-137, miR-138, miR-139, miR-140, miR-141, miR-142-3p, miR-143, miR-144, miR-145, miR-146, miR-147, miR-148a, miR-148b, miR-149, miR-150, miR-151, miR-152, miR-153, miR-154*, miR-154, miR-155, miR-181a, miR-181b, miR-181c, miR-182*, miR-182, miR-183, miR-184, miR-184, miR-185, miR-184, miR-18

185, miR-186, miR-187, miR-188, miR-189, miR-190, miR-191, miR-192, miR-193, miR-194, miR-195, miR-196a, miR-196b, miR-197, miR-198, miR-199a*, miR-199a, miR-199b, miR-200a, miR-200b, miR-200c, miR-202, miR-203, miR-204, miR-205, miR-206, miR-208, miR-210, miR-211, miR-212, miR-213, miR-213, miR-214, miR-215, miR-216, miR-217, miR-218, miR-220, miR-221, miR-222, miR-223, miR-224, miR-296, miR-299, miR-301, miR-302a*, miR-302a, miR-302b*, miR-302b, miR-302d, miR-302c*, miR-302c, miR-320, miR-323, miR-324-3p, miR-324-5p, miR-325, miR-326, miR-328, miR-330, miR-331, miR-337, miR-338, miR-339, miR-340, miR-342, miR-345, miR-346, miR-363, miR-367, miR-368, miR-370, miR-371, miR-372, miR-373*, miR-373, miR-374, miR-375, miR-376b, miR-378, miR-379, miR-380-5p, miR-380-3p, miR-381, miR-382, miR-383, miR-410, miR-412, miR-422a, miR-422b, miR-423, miR-424, miR-425, miR-429, miR-431, miR-448, miR-449, miR-450, miR-451, let7a, let7b, let7c, let7d, let7e, let7f, let7g, let7i, miR-376a, and miR-377. The sequences of the above-mentioned microRNA targets are provided in "the miRBase sequence database" as described in Griffith-Jones et al. (2004), Nucleic Acids Research 32:D109-D111, and Griffith-Jones et al. (2006), Nucleic Acids Research 34: D140-D144, which is publically accessible on the World Wide Web at the Wellcome Trust Sanger Institute website at http://microrna.sanger.ac.uk/sequences/.

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Exemplary primers for use in accordance with this embodiment of the kit are provided in TABLE 1, TABLE 2 and TABLE 6 below.

In another embodiment, the kit comprises one or more primer sets capable of detecting at least one or more of the following human microRNA target templates: miR-1, miR-7, miR-10b, miR-26a, miR-26b, miR-29a, miR-30e-3p, miR-95, miR-107, miR-141, miR-143, miR-154*, miR-154, miR-155, miR-181a, miR-181b, miR-181c, miR-190, miR-193, miR-194, miR-195, miR-202, miR-206, miR-208, miR-212, miR-221, miR-222, miR-224, miR-296, miR-299, miR-302c*, miR-302c, miR-320, miR-339, miR363, miR-376b, miR379, miR410, miR412, miR424, miR429, miR431, miR449, miR451, let7a, let7b, let7c, let7d, let7e, let7f, let7g, and let7i. Exemplary primers for use in accordance with this embodiment of the kit are provided in TABLE 1, TABLE 2 and TABLE 6 below.

In another embodiment, the kit comprises at least one oligonucleotide primer selected from the group consisting of SEQ ID NO: 2 to SEQ ID NO: 493, as shown in TABLE 1, TABLE 2, TABLE 6 and TABLE 7.

In another embodiment, the kit comprises at least one oligonucleotide primer selected from the group consisting of SEQ ID NO: 47, 48, 49, 50, 55, 56, 81, 82, 83, 84, 91, 92, 103, 104, 123, 124, 145, 146, 193, 194, 197, 198, 221, 222, 223, 224, 225, 226, 227, 228, 229, 230, 239, 240, 247, 248, 253, 254, 255, 256, 257, 258, 277, 278, 285, 286, 287, 288, 293, 294, 301, 302, 309, 310, 311, 312, 315, 316, 317, 318, 319, 320, 333, 334, 335, 336, 337, 338, 359, 360, 369, 370, 389, 390, 393, 394, 405, 406, 407, 408, 415, 416, 419, 420, 421, 422, 425, 426, 429, 430, 431, 432, 433, 434, 435, 436, 437, 438, 439, 440, 441, 442, 443, 444, 461 and 462, as shown in TABLE 6.

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A kit of the invention can also provide reagents for primer extension and amplification reactions. For example, in some embodiments, the kit may further include one or more of the following components: a reverse transcriptase enzyme, a DNA polymerase enzyme, a Tris buffer, a potassium salt (e.g., potassium chloride), a magnesium salt (e.g., magnesium chloride), a reducing agent (e.g., dithiothreitol), and deoxynucleoside triphosphates (dNTPs).

In various embodiments, the kit may include a detection reagent such as SYBR green dye or BEBO dye that preferentially or exclusively binds to double stranded DNA during a PCR amplification step. In other embodiments, the kit may include a forward and/or reverse primer that includes a fluorophore and quencher to measure the amount of the PCR amplification products.

The kit optionally includes instructions for using the kit in the detection and quantitation of one or more mammalian microRNA targets. The kit can also be optionally provided in a suitable housing that is preferably useful for robotic handling in a high throughput manner.

The following examples merely illustrate the best mode now contemplated for practicing the invention, but should not be construed to limit the invention.

EXAMPLE 1

This Example describes a representative method of the invention for producing DNA molecules from microRNA target molecules.

Primer extension was conducted as follows (using InVitrogen SuperScript III[®] reverse transcriptase and following the guidelines that were provided with the enzyme). The following reaction mixture was prepared on ice:

1 μl of 10 mM dNTPs

1 μl of 2 μM extension primer

1- 5 μl of target template
4 μl of "5X cDNA buffer"
1 μl of 0.1 M DTT
1 μl of RNAse OUT
1 μl of SuperScript III[®] enzyme

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The mixture was incubated at 50°C for 30 minutes, then 85°C for 5 minutes, then cooled to room temperature and diluted 10-fold with TE (10 mM Tris, pH 7.6, 0.1 mM EDTA).

Real-time PCR was conducted using an ABI 7900 HTS detection system (Applied Biosystems, Foster City, California, U.S.A.) by monitoring SYBR® green fluorescence of double-stranded PCR amplicons as a function of PCR cycle number. A typical 10 μl PCR reaction mixture contained:

5 μl of 2X SYBR® green master mix (ABI)

0.8 μl of 10 μM universal forward primer

0.8 μl of 10 μM reverse primer

1.4 µl of water

water to 20 µl

2.0 µl of target template (10-fold diluted RT reaction).

The reaction was monitored through 40 cycles of standard "two cycle" PCR $(95^{\circ}\text{C} - 15 \text{ sec}, 60^{\circ}\text{C} - 60 \text{ sec})$ and the fluorescence of the PCR products was measured.

The foregoing method was successfully used in eleven primer extension PCR assays for quantitation of endogenous microRNAs present in a sample of total RNA. The DNA sequences of the extension primers, the universal forward primer sequence, and the LNA substituted reverse primers, used in these 11 assays are shown in TABLE 1.

25 TABLE 1

Target microRNA	Primer	Primer Name	DNA sequence (5' to 3')	SEQ ID
	number			NO
gene-specific extens	sion primer:	s ¹		
humanb let7a	357	let7aP4	CATGATCAGCTGGGCCAAGAAACTATACAACCT	2
human miR-1	337	miR1P5	CATGATCAGCTGGGCCAAGATACATACTTCT	3
human miR-15a	344	miR15aP3	CATGATCAGCTGGGCCAAGACAAAACCATTATG	4
human miR-16	351	miR16P2	CATGATCAGCTGGGCCAAGACGCCAATA <u>TTTACGT</u>	5

Target microRNA	Primer	Primer Name	DNA sequence (5' to 3')	SEQ ID
	number			NO
human miR-21	342	miR21P6	CATGATCAGCTGGGCCAAGATCAACATCAGT	6
human miR-24	350	miR24P5	CATGATCAGCTGGGCCAAGACTGTTCCTGCTG	7
human miR-122	222	122-E5F	CATGATCAGCTGGGCCAAGAACAAACACCA <u>TTGTCA</u>	8
human miR-124	226	124-E5F	CATGATCAGCTGGGCCAAGATGGCATTCACCGCGTG	9
human miR-143	362	miR143P5	CATGATCAGCTGGGCCAAGATGAGCTA <u>CAGTG</u>	10
human miR-145	305	miR145P2	CATGATCAGCTGGGCCAAGAAAGGGATTCCTGGGAA	11
human miR-155	367	miR155P3	CATGATCAGCTGGGCCAAGACCCCTATCACGAT	12
¹ - Universal forwar	d primer bi	nding sites are sh	own in italics. The overlap with the RNA-specific reverse	
primers are underlin	ned.			
universal forward p	rimer			
*	230	E5F	CATGATCAGCTGGGCCAAGA	13
RNA species-specif	ic reverse p	orimers ²		
human let7a	290	miRlet7a-	TG+AGGT+AGT <u>AGGTTG</u>	14
		1,2,3R		
human miR-1	285	miR1-1,2R	TG+GAA+TG+TAA <u>AGAAGTA</u>	15
human miR-15a	287	miR15aR	TAG+CAG+CACATAATG	16
human miR-16	289	miR16-1,2R	T+AGC+AGC <u>ACGTAAA</u>	17
human miR-21	286	miR21R	T+AG+CT+TATCAG <u>ACTGAT</u>	18
human miR-24	288	miR24-1,2R	TGG+CTCAGTT <u>CAGC</u>	19
human miR-122	234	122LNAR	T+G+GAG+TG <u>TGACAA</u>	20
human miR-124	235	124LNAR	T+TAA+GG <u>CACGCG</u>	21
human miR-143	291	miR143R	TG+AGA+TGAAG <u>CACTG</u>	22
human miR-145	314	miR145R2	GT+CCAGTT <u>TTCCCA</u>	23
human miR-155	293	miR155R	T+TAA+TG+CTA <u>ATCGTGA</u>	24
			on of overlap of the reverse primers with the corresponding	
extension primers ar			.r k	
panarolo ul				

The assay was capable of detecting microRNA in a concentration range of from 2 nM to 20 fM. The assays were linear at least up to a concentration of 2 nM of synthetic microRNA (>1,000,000 copies/cell).

EXAMPLE 2

This Example describes the evaluation of the minimum sequence requirements for efficient primer-extension mediated cDNA synthesis using a series of extension primers for microRNA assays having gene specific regions that range in length from 12 to 3 base pairs.

Primer Extension Reactions: Primer extension was conducted using the target molecules miR-195 and miR-215 as follows. The target templates miR-195 and miR-215 were diluted to 1nM RNA (100,000 copies/cell) in TE zero plus 100ng/ll total yeast RNA. A no template control (NTC) was prepared with TE zero plus 100ng/ll total yeast RNA.

The reverse transcriptase reactions were carried out as follows (using InVitrogen SuperScript III[®] reverse transcriptase and following the guidelines that were provided with the enzyme) using a series of extension primers for miR-195 (SEQ ID NO: 25-34) and a series of extension primers for miR-215 (SEQ ID NO: 35-44) the sequences of which are shown below in TABLE 2.

The following reaction mixtures were prepared on ice:

Set 1: No Template Control

37.5µl water

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12.5µl of 10mM dNTPs

12.5µl 0.1 mM DTT

20 50μl of "5X cDNA buffer"

12.5µl RNAse OUT

12.5 µl Superscript III® reverse transcriptase enzyme

12.5 µl lugul Hela cell total RNA (Ambion)

plus 50µl of 2µM extension primer

25 plus 50µl TEzero + yeast RNA

Set 2: Spike-in Template

37.5 µl water

12.5µl of 10mM dNTPs

30 12.5µl 0.1 mM DTT

50µl of "5X cDNA buffer"

12.5µl RNAse OUT

12.5µl Superscript III® reverse transcriptase enzyme (InVitrogen)

12.5 µl µg/u Hela cell total RNA (Ambion)

plus 50µl of 2µM extension primer

plus 50µl 1 nM RNA target template (miR-195 or miR-215) serially diluted in 10-5 fold increments

The reactions were incubated at 50°C for 30 minutes, then 85°C for 5 minutes, and cooled to 4°C and diluted 10-fold with TE (10mM Tris, pH 7.6, 0.1 mM EDTA).

Quantitative Real-Time PCR reactions: Following reverse transcription, quadruplicate measurements of cDNA were made by quantitative real-time (qPCR) using an ABI 7900 HTS detection system (Applied Biosystems, Foster City, California, U.S.A.) by monitoring SYBR® green fluorescence of double-stranded PCR amplicons as a function of PCR cycle number. The following reaction mixture was prepared:

5µl of 2X SYBR green master mix (ABI)

0.8µl of 10µM universal forward primer (SEQ ID NO: 13)

0.8μl of 10μM reverse primer (miR-195RP:SEQ ID NO: 45 or miR215RP: SEQ ID NO: 46)

1.4µl of water

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2.0µl of target template (10-fold diluted miR-195 or miR-215 RT reaction)

Quantitative real-time PCR was performed for each sample in quadruplicate, using the manufacturer's recommended conditions. The reactions were monitored through 40 cycles of standard "two cycle" PCR (95°C – 15 sec, 60°C – 60 sec) and the fluorescence of the PCR products were measured and disassociation curves were generated. The DNA sequences of the extension primers, the universal forward primer sequence, and the LNA substituted reverse primers, used in the miR-195 and miR-215 assays are shown below in TABLE 2. The assay results for miR-195 are shown below in TABLE 3 and the assay results for miR-215 are shown below in TABLE 4.

TABLE 2

IAD	LE 2	r		
Target microRNA	Primer number	Primer Name	DNA sequence (5' to 3')	SEQ ID NO:
gene-specifi	c extensio	on primers ¹		
miR-195	646	mir195- GS1	CATGATCAGCTGGGCCAAGAGCCAATATTTCT	25
miR-195	647	mir195- GS2	CATGATCAGCTGGGCCAAGAGCCAATATTTC	26
miR-195	648	mir195- GS3	CATGATCAGCTGGGCCAAGAGCCAATATTT	27
miR-195	649	mir195- GS4	CATGATCAGCTGGGCCAAGAGCCAATATT	28
miR-195	650	mir195- GS5	CATGATCAGCTGGGCCAAGAGCCAATAT	29
miR-195	651	mir195- GS6	CATGATCAGCTGGGCCAAGAGCCAATA	30
miR-195	652	mir195- GS7	CATGATCAGCTGGGCCAAGAGCCAAT	31
miR-195	653	mir195- GS8	CATGATCAGCTGGGCCAAGAGCCAA	32
miR-195	654	mir195- GS9	CATGATCAGCTGGGCCAAGAGCCA	33
miR-195	655	mir195- GS10	CATGATCAGCTGGGCCAAGAGCC	34
miR-215	656	mir215- GS1	CATGATCAGCTGGGCCAAGAGTCTGTCAATTC	35
miR-215	657	mir215- GS2	CATGATCAGCTGGGCCAAGAGTCTGTCAATT	36
miR-215	658	mir215- GS3	CATGATCAGCTGGGCCAAGAGTCTGTCAAT	37
miR-215	659	mir215- GS4	CATGATCAGCTGGGCCAAGAGTCTGTCAA	38

Target microRNA	Primer number	Primer Name	DNA sequence (5' to 3')	SEQ ID NO:
miR-215	660	mir215- GS5	CATGATCAGCTGGGCCAAGAGTCTGTCA	39
miR-215	661	mir215- GS6	CATGATCAGCTGGGCCAAGAGTCTGTC	40
miR-215	662	mir215- GS7	<i>CATGATCAGCTGGGCCAAGA</i> GTCTGT	41
miR-215	663	mir215- GS8	CATGATCAGCTGGGCCAAGAGTCTG	42
miR-215	664	mir215- GS9	<i>CATGATCAGCTGGGCCAAGA</i> GTCT	43
miR-215	665	mir215- GS10	<i>CATGATCAGCTGGGCCAAGA</i> GTC	44
¹ - Universa	l forward	primer bindi	ng sites are shown in italics.	
RNA specie	s-specific	reverse prin	ners ²	
miR-195	442	mir195RP	T+AGC+AGCACAGAAAT	45
miR-215	446	mir215RP	A+T+GA+CCTATGAATTG	46
² - The "+" s	symbol pro	ecedes the Ll	NA molecules.	

Results:

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The sensitivity of each assay was measured by the cycle threshold (Ct) value which is defined as the cycle count at which fluorescence was detected in an assay containing microRNA target template. The lower this Ct value (e.g. the fewer number of cycles), the more sensitive was the assay. For microRNA samples, it was generally observed that while samples that contain template and no template controls both eventually cross the detection threshold, the samples with template do so at a much lower cycle number. The Δ Ct value is the difference between the number of cycles (Ct) between template containing samples and no template controls, and serves as a measure of the dynamic range of the assay. Assays with a high dynamic range allow measurements of very low microRNA copy numbers. Accordingly, desirable

characteristics of a microRNA detection assay include high sensitivity (low Ct value) and broad dynamic range (Δ Ct \geq 12) between the signal of a sample containing target template and a no template background control sample.

The results of the miR195 and miR215 assays using extension primers having a gene specific portion ranging in size from 12 nucleotides to 3 nucleotides are shown below in TABLE 3 and TABLE 4, respectively. The results of these experiments unexpectedly demonstrate that gene-specific priming sequences as short as 3 nucleotides exhibit template specific priming. For both the miR-195 assay sets (shown in TABLE 3) and the miR-215 assay sets (shown in TABLE 4), the results demonstrate that the dynamic range (Δ Ct) for both sets of assays are fairly consistent for extension primers having gene specific regions that are greater or equal to 8 nucleotides in length. The dynamic range of the assay (Δ Ct) begins to decrease for extension primers having gene specific regions below 8 nucleotides, with a reduction in assay specificity below 7 nucleotides in the miR-195 assays, and below 6 nucleotides in the miR-215 assays. A melting point analysis of the miR-215 samples demonstrated that even at 3 nucleotides, there is specific PCR product present in the plus template samples (data not shown). Taken together, these data demonstrate that the gene specific region of extension primers is ideally \geq 8 nucleotides, but can be as short as 3 nucleotides in length.

TABLE 3: miR195 Assay Results

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GS Primer Length	Ct: No Template	Ct: Plus Template	Δ Ct
12	34.83	20.00	14.82
12	34.19	19.9	14.3
11	40.0	19.8	20.2
10	36.45	21.2	15.2
9	36.40	22.2	14.2
8	40.0	23.73	16.27
7	36.70	25.96	10.73
6	30.95	26.58	4.37

GS Primer Length	Ct: No Template Control	Ct: Plus Template	Δ Ct
5	30.98	31.71	-0.732
4	32.92	33.28	-0.364
3	35.98	35.38	-0.605

Ct=the cycle count where the fluorescence exceeds the threshold of detection. Δ Ct = the difference between the Ct value with template and no template.

TABLE 4: miR215 Assay Results

GS Primer Length	Ct: No Template Control	Ct: Plus Template	Δ Ct
12	33.4	13.57	19.83
12	33.93	14.15	19.77
11	35.51	15.76	19.75
10	35.33	15.49	19.84
9	36.02	16.84	19.18
8	35.79	17.07	18.72
7	32.29	17.58	14.71
6	34.38	20.62	13.75
5	34.41	28.65	5.75
4	36.36	33.92	2.44
3	35.09	33.38	1.70

Ct=the cycle count where the fluorescence exceeds the threshold of detection. Δ Ct = the difference between the Ct value with template and no template.

EXAMPLE 3

This Example describes assays and primer sets designed for quantitative analysis of human microRNA expression patterns.

Primer Design:

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microRNA target templates: the sequence of the target templates as described herein are publically available accessible on the World Wide Web at the Wellcome Trust Sanger Institute website in the "miRBase sequence database" as described in Griffith-Jones et al. (2004), Nucleic Acids Research 32:D109-D111 and and Griffith-Jones et al. (2006), Nucleic Acids Research 34: D140-D144.

Extension primers: gene specific primers for primer extension of a microRNA to form a cDNA followed by quantitative PCR (qPCR) amplification were designed to (1) convert the RNA template into cDNA; (2) to introduce a "universal" PCR binding site (SEQ ID NO:1) to one end of the cDNA molecule; and (3) to extend the length of the cDNA to facilitate subsequent monitoring by qPCR.

Reverse primers: unmodified reverse primers and locked nucleic acid (LNA) containing reverse primers (RP) were designed to quantify the primer-extended, full length cDNA in combination with a generic universal forward primer (SEQ ID NO:13). For the locked nucleic acid containing reverse primers, two or three LNA modified bases were substituted within the first 8 nucleotides from the 5' end of the reverse primer oligonucleotide, as shown below in the exemplary reverse primer sequences provided in TABLE 6. The LNA base substitutions were selected to raise the predicted Tm of the primer by the highest amount, and the final predicted Tm of the selected primers were specified to be preferably less than or equal to 55°C.

An example describing an assay utilizing an exemplary set of primers the detection of miR-95 and miR-424 is described below.

<u>Primer Extension Reactions</u>: primer extension was conducted using DNA templates corresponding to miR-95 and miR-424 as follows. The DNA templates were diluted to 0 nM, 1 nM, 100 pM, 10 pM and 1 pM dilutions in TE zero (10 mM Tris pH7.6, 0.1 mM EDTA) plus 100ng/µl yeast total RNA (Ambion, Austin TX).

The reverse transcriptase reactions were carried out using the following primers:

<u>Extension primers:</u> (diluted to 500 nM)

miR-95GSP CATGATCAGCTGGGCCAAGATGCTCAATAA (SEQ ID NO:

123)

miR-424GSP CATGATCAGCTGGGCCAAGATTCAAAACAT (SEQ ID NO:415)

Reverse primers: (diluted to 10 mM).

miR-95_RP4 TT+CAAC+GGGTATTTATTGA (SEQ ID NO: 124)

miR-424RP2 C+AG+CAGCAATTCATGTTTT (SEQ ID NO: 416)

Reverse Transcription (per reaction):

2µl water

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2µl of "5X cDNA buffer" (InVitrogen, Carlsbad, CA)

10 0.5μl of 0.1 mM DTT (InVitrogen, Carlsbad, CA)

0.5 µl of 10 mM dNTPs (InVitrogen, Carlsbad, CA)

0.5 µl RNAse OUT (InVitrogen, Carlsbad, CA)

0.5µl Superscript III[®] reverse transcriptase enzyme (InVitrogen, Carlsbad, CA)

2µl of extension primer plus 2µl of template dilution.

The reactions were mixed and incubated at 50°C for 30 minutes, then 85°C for 5 minutes, and cooled to 4°C and diluted 10-fold with TE zero.

Quantitative Real-Time PCR Reactions: (per reaction)

5µl 2X SYBR mix (Applied Biosystems, Foster City, CA)

1.4µl water

0.8 µl universal primer (CATGATCAGCTGGGCCAAGA (SEQ ID NO: 13))

2.0µl of diluted reverse transcription (RT) product from above.

Quantitative real-time PCR was performed for each sample in quadruplicate, using the manufacturer's recommended conditions. The reactions were monitored through 40 cycles of standard "two cycle" PCR (95°C – 15 sec, 60°C – 60 sec) and the fluorescence of the PCR products were measured and disassociation curves were generated. The DNA sequences of the extension primers, the universal forward primer sequence, and the LNA substituted reverse primers, used in the representative miR-95 and miR-424 assays as well as primer sets for 212 different human microRNA templates are shown below in TABLE 6. Primer sets for assays requiring extensive testing and design modification to achieve a sensitive assay with a high dynamic range are indicated in TABLE 6 with the symbol # following the primer name.

Results:

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TABLE 5 shows the Ct values (averaged from four samples) from the miR-95 and miR-424 assays, which are plotted in the graph shown in FIGURE 2. The results of these assays are provided as representative examples in order to explain the significance of the assay parameters shown in TABLE 6 designated as slope (column 6), intercept (column 7) and background (column 8).

As shown in TABLE 5, the Ct value for each template at various concentrations is provided. The Ct values (x-axis) are plotted as a function of template concentration (y-axis) to generate a standard curve for each assay, as shown in FIGURE 2. The slope and intercept define the assay measurement characteristics that permit an estimation of number of copies/cell for each microRNA. For example, when the Ct values for 50 µg total RNA input for the miR-95 assay are plotted, a standard curve is generated with a slope and intercept of -.03569 and 9.655, respectively. When these standard curve parameters are applied to the Ct of an unknown sample (x), they yield log 10 (copies/20pg total RNA) (y). Because the average cell yields 20 pg of total RNA, these measurements equate to copies of microRNA/cell. The background provides an estimate of the minimum copy number that can be measured in a sample and is computed by inserting the no template control (NTC) value into this equation. In this example, as shown in TABLE 6, miR-95 yields a background of 1.68 copies/20 pg at 50 µg of RNA input.

As further shown in TABLE 6, reverse primers that do not contain LNA may also be used in accordance with the methods of the invention. See, e.g. SEQ ID NO: 494-499. The sensitivity and dynamic range of the assays using non-LNA containing reverse primers SEQ ID NO: 494-499, yielded similar results to the corresponding assays using LNA-containing reverse primers.

TABLE 5

Ct Values (avera	aged from four	samples)				
Template concentration	10 nM	1 nM	0.1 nM	0.01 nM	0.001 nM	NTC
copies/20 pg RNA (50 μg input)	500,000	50,000	5000	500	50	

Ct Values (avera	aged from four s	samples)				
Template concentration	10 nM	1 nM	0.1 nM	0.01 nM	0.001 nM	NTC
copies/20 pg RNA (5 µg input)	5,000,000	500,000	50,000	5000	500	
miR-95	11.71572163	14.17978	17.46353	19.97259	23.33171	27.44383
miR-424	10.47708975	12.76806	15.69251	18.53729	21.56897	23.2813
log10 (copies for 50 µg input)	5.698970004	4.69897	3.69897	2.69897	1.69897	

TABLE 6: Primers to detect human microRNA target templates

Human Target micro	Extension		Reverse				Backg	Background RNA input
RNA	Primer Name	Extension Primer Sequence	Name	Reverse Primer Sequence	Slope	Intercept	50ug	Śug
# denotes pi	rimers for assays th	# denotes primers for assays that required extensive testing and primer design modification to achieve optimal assay results including high sensitivity and high dynamic range.	to achieve optimal	assay results including high sensitiv	ity and high	ı dynamic range	വ്	
miR-1	miR1GSP10#	CATGATCAGCTGGGCCAAGATACATACTTC	miR-1RP#	T+G+GAA+TG+TAAAGAAGT	-0.2758	8.3225	2.44	24.36
		SEQ ID NO:47		SEQ ID NO:48				
miR-7	miR-7GSP10#	CATGATCAGCTGGGCCAAGACAACAAAATC	miR-7 RP6#	T+GGAA+GACTAGTGATTTT	-0.2982	10.435	11.70	116.99
:		SEQ ID NO:49	1	SEQ ID NO:50				
miR-9*	miR-9*GSP	CATGATCAGCTGGGCCAAGAACTTTCGGTT	miR-9*RP	TAAA+GCT+AGATAACCG	-0.2405	8.9145	3.71	37.15
		SEQ ID NO:51		SEQ ID NO:52				
miR-10a	miR-10aGSP	CATGATCAGCTGGGCCAAGACACAAATTCG	miR-10aRP	T+AC+CCTGTAGATCCG	-0.2755	926978	60.0	0.94
		SEQ ID NO:53		SEQ ID NO:54				
miR-10b	miR-	CATGATCAGCTGGGCCAAGAACAAATTCGGT	miR-	TA+CCC+TGT+AGAACCGA	-0.3505	8.7109	0.55	5.52
	10b_GSP11#	SEQ ID NO:55	10b_RP2#	SEQ ID NO:56				
miR-15a	miR-15aGSP	CATGATCAGCTGGGCCAAGACACAAACCAT	miR-15aRP	T+AG+CAGCACATAATG	-0.2831	8.4519	4.40	44.01
		SEQ ID NO:57		SEQ ID NO:58				
miR-15b	miR-15bGSP2	CATGATCAGCTGGGCCAAGATGTAAACCA	miR-15bRP	T+AG+CAGCACATCAT	-0.2903	8.4206	0.18	1.84
		SEQ ID NO:59		SEQ ID NO:60				
miR-16	miR-16GSP2	CATGATCAGCTGGGCCAAGACGCCAATAT	miR-16RP	T+AG+CAGCACGTAAA	-0.2542	9.3689	1.64	16.42
		SEQ ID NO:61		SEQ ID NO:62				
miR-17-	miR-17-3pGSP	CATGATCAGCTGGGCCAAGAACAAGTGCCT	miR-17-3pRP	A+CT+GCAGTGAAGGC	-0.2972	8.2625	1.08	10.78
3p		SEQ ID NO:63		SEQ ID NO:64				
miR-17-	miR-17-	CATGATCAGCTGGGCCAAGAACTACCTGC	miR-17-5pRP	C+AA+AGTGCTTACAGTG	-0.2956	7.9101	0.13	1.32
dc	5pGSP2	SEQ ID NO:65		SEQ ID NO:66				

RNA Primer Name Extens IIR-19a miR-19aGSP2 CATGATCAGCTGG IIR-19b SEQ ID NO:67 SEQ ID NO:67 miR-20 CATGATCAGCTGG SEQ ID NO:69 miR-20 miR-21GSP2 CATGATCAGCTGG miR-21 miR-21GSP2 CATGATCAGCTGG miR-23 miR-21GSP2 CATGATCAGCTGG SEQ ID NO:73 SEQ ID NO:73 miR-23 miR-236GSP CATGATCAGCTGG miR-25 miR-25GSP CATGATCAGCTGG miR-26 SEQ ID NO:77 SEQ ID NO:79 miR-26 SEQ ID NO:79 SEQ ID NO:79 miR-26 SEQ ID NO:79 SEQ ID NO:79 miR-26 SEQ ID NO:81 SEQ ID NO:81 miR-26 SEQ ID NO:81 SEQ ID NO:83 miR-27a SEQ ID NO:83 SEQ ID NO:83 miR-27b SEQ ID NO:85 SEQ ID NO:85 miR-27b SEQ ID NO:85 SEQ ID NO:85		Reverse Primer				Background RNA input	ound
miR-19aGSP2 miR-20GSP3 miR-21GSP2 miR-23aGSP miR-23aGSP miR-25GSP miR-25GSP miR-26aGSP9# miR-27aGSP	Extension Primer Sequence	Name	Reverse Primer Sequence	Slope	Intercept	20ug	Sug
miR-20GSP3 miR-23aGSP miR-23bGSP miR-25GSP miR-26bGSP9# miR-26bGSP9# miR-27aGSP	CTGGGCCAAGATCAGTTTTG	miR-19aRP	TG+TG+CAAATCTATGC	-0.2984	9.461	0.02	0.23
miR-20GSP3 miR-23aGSP miR-23bGSP miR-25GSP miR-26aGSP9# miR-26bGSP9# miR-27aGSP) NO:6/		SEQ ID NO:68				
miR-21GSP2 miR-23aGSP miR-23bGSP miR-26aGSP9# miR-26aGSP9# miR-26aGSP9#	CATGATCAGCTGGGCCAAGATCAGTTTTGC	miR-19bRP	TG+TG+CAAATCCATG	-0.294	8.1434	2.26	22.55
miR-20GSP3 miR-23aGSP miR-23bGSP miR-25GSP miR-26aGSP9# miR-26bGSP9# miR-27aGSP	9 NO:69		SEQ ID NO:70				
miR-23aGSP miR-23bGSP miR-25GSP miR-26aGSP9# miR-26bGSP9# miR-27aGSP	CATGATCAGCTGGCCAAGACTACCTGC	miR-20RP	T+AA+AGTGCTTATAGTGCA	-0.2979	7.9929	0.16	1.60
miR-21GSP2 miR-23aGSP miR-23bGSP miR-25GSP miR-26aGSP9# miR-26aGSP9# miR-27aGSP) NO:71		SEQ ID NO:72				
miR-23aGSP miR-23bGSP miR-26aGSP9# miR-26bGSP9# miR-27aGSP	CATGATCAGCTGGGCCAAGATCAACATCA	miR-21RP	T+AG+CTTATCAGACTGATG	-0.2849	8.1624	1.80	17.99
miR-23aGSP miR-23bGSP miR-26aGSP9# miR-26bGSP9# miR-27aGSP) NO: 73		SEQ ID NO:74				
miR-23bGSP miR-26aGSP9# miR-26bGSP9# miR-27aGSP	CATGATCAGCTGGGCCAAGAGGAAATCCCT	miR-23aRP	A+TC+ACATTGCCAGG	-0.3172	9.4253	2.41	24.08
miR-23bGSP miR-26aGSP9# miR-26bGSP9# miR-27aGSP	NO:75		SEQ ID NO:76				
miR-25GSP miR-26aGSP9# miR-27aGSP9 miR-27aGSP	CATGATCAGCTGGGCCAAGAGGTAATCCCT	miR-23bRP	A+TC+ACATTGCCAGG	-0.2944	9.0985	5.39	53.85
miR-25GSP miR-26aGSP9# miR-26bGSP9# miR-27aGSP	71.ON C		SEQ ID NO:78				:
miR-26aGSP9# miR-26bGSP9# miR-27aGSP	CATGATCAGCTGGGCCAAGATCAGACCGAG	miR-25RP	C+AT+TGCACTTGTCTC	-0.3009	8.2482	1.52	15.19
miR-26aGSP9# miR-26bGSP9# miR-27aGSP	0 NO:79		SEQ ID NO:80				
miR-26bGSP9# miR-27aGSP miR-27bGSP	CATGATCAGCTGGGCCAAGAGCCTATCCT	miR-	TT+CA+AGIAATCCAGGAT	-0.2807	8.558	0.26	2.56
miR-26bGSP9# miR-27aGSP miR-27bGSP		26aRP2#	SEQ ID NO:82				
miR-27aGSP miR-27bGSP	CATGATCAGCTGGGCCAAGAAACCTATCC	miR-	TT+CA+AGT+AATTCAGGAT	-0.2831	8.7885	0.37	3.67
miR-27aGSP miR-27bGSP		26bRP2#	SEQ ID NO:84				
miR-27bGSP	CATGATCAGCTGGGCCAAGAGCGGAACTTA	miR-27aRP	TT+CA+CAGTGGCTAA	-0.2765	9.5239	5.15	51.51
miR-27bGSP	O NO:85		SEQ ID NO:86				
	CATGATCAGCTGGGCCAAGAGCAGAACTTA	miR-27bRP	TT+CA+CAGTGGCTAA	-0.28	9.5483	5.97	59.71
SEQ ID NO:87	NO:87		SEQ ID NO:88				
miR-28 miR-28GSP CATGATCAGCTGG	CATGATCAGCTGGGCCAAGACTCAATAGAC	miR-28RP	A+AG+GAGCTCACAGT	-0.3226	10.01	7.19	71.87
SEQ ID NO:89	O NO:89		SEQ ID NO:90				

PCT/US2006/002591

WO 2006/081284

Human Target micro RNA	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Slope	Intercept	Background RNA input 50ug 5ug	round input Sug
iR-29a	miR-29aGSP8#	CATGATCAGCTGGGCCAAGAAACCGATT SEQ ID NO:91	miR- 29aRP2#	T+AG+CACCATCTGAAAT SEQ ID NO:92	-0.29	8.8731	0.04	0.38
iR-29b	miR-29bGSP2	CATGATCAGCTGGGCCAAGAAACACTGAT SEQ ID NO:93	miR-29bRP2	T+AG+CACCATTTGAAATCA G SEQ ID NO:94	-0.3162	9.6276	3.56	35.57
miR-30a- 5p	miR-30a- 5pGSP	CATGATCAGCTGGGCCAAGACTTCCAGTCG SEQ ID NO:95	miR-30a- 5pRP	T+GT+AAACATCCTCGAC SEQ ID NO:96	-0.2772	9.0694	1.92	19.16
miR-30b	miR-30bGSP	CATGATCAGCTGGGCCAAGAAGCTGAGTGT SEQ ID NO:97	miR-30bRP	TGT+AAA+CATCCTACACT SEQ ID NO:98	-0.2621	8.5974	0.11	1.13
miR-30c	miR-30cGSP	CATGATCAGCTGGGCCAAGAGCTGAGAGTG SEQ ID NO:99	miR-30cRP	TGT+AAA+CATCCTACACT SEQ ID NO:100	-0.2703	8.699	0.15	1.48
miR-30d	miR-30dGSP	CATGATCAGCTGGGCCAAGACTTCCAGTCG SEQ ID NO:101	miR-30dRP	T+GTAAA+CATCCCCG SEQ ID NO:102	-0.2506	9.3875	0.23	2.31
miR-30e- 3p	miR-30e- 3pGSP9#	CATGATCAGCTGGGCCAAGAGCTGTAAAC SEQ ID NO:103	miR-30e- 3pRP5#	CTTT+CAGT+CGGATGTTT SEQ ID NO:104	-0.325	11.144	6.37	63.70
miR-30e- 5p	miR-30e- 5pGSP	CATGATCAGCTGGGCCAAGATCCAGTCAAG SEQ ID NO:105	miR-30e- 5pRP	TG+TAAA+CATCCTTGAC SEQ ID NO:106	-0.2732	8.1604	8.50	85.03
miR-31	miR-31GSP	CATGATCAGCTGGGCCAAGACAGCTATGCC SEQ ID NO:107	miR-31RP	G+GC+AAGATGCTGGC SEQ ID NO:108	-0.3068	8.2605	3.74	37.43
miR-32	miR-32GSP	CATGATCAGCTGGGCCAAGAGCAACTTAGT SEQ ID NO:109	miR-32RP	TATTG+CA+CATTACTAAG SEQ ID NO:110	-0.2785	8.9581	0.39	3.93
miR-33	miR-33GSP2	CATGATCAGCTGGGCCAAGACAATGCAAC SEQ ID NO:111	miR-33RP	G+TG+CATTGTAGTTGC SEQ ID NO:112	-0.3031	8.42	2.81	28.14
miR-34a	miR-34aGSP	CATGATCAGCTGGGCCAAGAAACAACCAGC SEQ ID NO:113	miR-34aRP	T+GG+CAGTGTCTTAG SEQ ID NO:114	-0.3062	9.1522	2.40	23.99

Human Target micro RNA	Extension Primer Name	Petancian Deimor Commen	Reverse Primer	Daving Dimos Contract	Slone	Tutomote	Backgr RNA ii	Background RNA input
iiR-34b	miR-34bGSP	CATGATCAGCTGGGCCAAGACAATCAGCTA	miR-34bRP	TA+GG+CAGTGTCATT	-0.3208	9.054	0.04	0.37
		SEQ ID NO:115		SEQ ID NO:116				
iR-34c	miR-34cGSP	CATGATCAGCTGGGCCAAGAGCAATCAGCT	miR-34cRP	A+GG+CAGTGTAGTTA	-0.2995	10.14	1.08	10.83
		SEQ ID NO:117		SEQ ID NO:118				
miR-92	miR-92GSP	CATGATCAGCTGGGCCAAGACAGGCCGGGA	miR-92RP	T+AT+TGCACTTGTCCC	-0.3012	8069'8	8.92	89.17
		SEQ ID NO:119		SEQ ID NO:120				
miR-93	miR-93GSP	CATGATCAGCTGGGCCAAGACTACCTGCAC	miR-93RP	AA+AG+TGCTGTTCGT	-0.3025	7.9933	4.63	46.30
		SEQ ID NO:121		SEQ ID NO:122				
miR-95	miR-95GSP#	CATGATCAGCTGGGCCAAGATGCTCAATAA	miR-	TT+CAAC+GGGTATTTATTG	-0.3436	559.6	1.68	16.80
		SEQ ID NO:123	95_RP4#	A				
				SEQ ID NO:124				
miR-96	miR-96GSP	CATGATCAGCTGGGCCAAGAGCAAAAATGT	miR-96RP	T+TT+GGCACTAGCAC	-0.2968	9.2611	0.00	0.05
		SEQ ID NO:125		SEQ ID NO:126				
miR-98	miR-98GSP	CATGATCAGCTGGGCCAAGAAACAATACAA	miR-98RP	TGA+GGT+AGTAAGTTG	<i>-</i> 0.2797	9.5654	1.05	10.48
		SEQ ID NO:127		SEQ ID NO:128				
miR-99a	miR-99aGSP	CATGATCAGCTGGGCCAAGACACAAGATCG	miR-99aRP	A+AC+CCGTAGATCCG	-0.2768	8.781	0.21	2.08
		SEQ ID NO:129		SEQ ID NO:130			_	
miR-99b	miR-99bGSP	CATGATCAGCTGGGCCAAGACGCAAGGTCG	miR-99bRP	C+AC+CCGTAGAACCG	-0.2747	7.9855	0.25	2.53
		SEQ ID NO:131		SEQ ID NO:132				
miR-100	miR-100GSP	CATGATCAGCTGGGCCAAGACACAAGTTCG	miR-100RP	A+AC+CCGTAGATCCG	-0.2902	699.8	0.04	0.35
		SEQ ID NO:133		SEQ ID NO:134				
miR-101	miR-101GSP	CATGATCAGCTGGGCCAAGACTTCAGTTAT	miR-101RP	TA+CAG+TACTGTGATAACT	-0.3023	8.2976	0.46	4.63
		SEQ ID NO:135		SEQ ID NO:136				
miR-103	miR-103GSP	CATGATCAGCTGGGCCAAGATCATAGCCCT	miR-103RP	A+GC+AGCATTGTACA	-0.3107	8.5776	0.02	0.21
		SEQ ID NO:137		SEQ ID NO:138				

Human								
Target micro	Extension		Reverse Primer	-			Background RNA input	round input
RNA	Primer Name	Extension Primer Sequence	Name	Reverse Primer Sequence	Slope	Intercept	20ug	Sug
iR-105	miR-105GSP	CATGATCAGCTGGGCCAAGAACAGGAGTCT	miR-105RP	T+CAAA+TGCTCAGACT	-0.2667	8.9832	0.93	9.28
		SEQ ID NO:139		SEQ ID NO:140				
iR-106a	miR-106aGSP	CATGATCAGCTGGGCCAAGAGCTACCTGCA	miR-106aRP	AAA+AG+TGCTTACAGTG	-0.3107	8.358	0.03	0.31
		SEQ ID NO:141		SEQ ID NO:142				
miR-106b	miR-106bGSP	CATGATCAGCTGGGCCAAGAATCTGCACTG	miR-106bRP	T+AAAG+TGCTGACAGT	-0.2978	8.7838	0.10	1.04
		SEQ ID NO:143		SEQ ID NO:144				
miR-107	miR-107GSP8#	CATGATCAGCTGGGCCAAGATGATAGCC	miR-	A+GC+AGCATTGTACAG	-0.304	9.1666	0.34	3.41
		SEQ ID NO:145	107RP2#	SEQ ID NO:146				
miR-122a	miR-122aGSP	CATGATCAGCTGGGCCAAGAACAACACCA	miR-122aRP	T+GG+AGTGTGACAAT	-0.3016	8.1479	90.0	0.58
		SEQ ID NO:147		SEQ ID NO:148				
miR-124a	miR-124aGSP	CATGATCAGCTGGGCCAAGATGGCATTCAC	miR-124aRP	T+TA+AGGCACGCGGT	-0.3013	9069.8	0.56	5.63
		SEQ ID NO:149		SEQ ID NO:150				
miR-125a	miR-125aGSP	CATGATCAGCTGGGCCAAGACACAGGTTAA	miR-125aRP	T+CC+CTGAGACCCTT	-0.2938	8.6754	60.0	0.91
		SEQ ID NO:151		SEQ ID NO:152				
miR-125b	miR-125bGSP	CATGATCAGCTGGGCCAAGATCACAAGTTA	miR-125bRP	T+CC+CTGAGACCCTA	-0.283	8.1251	0.20	1.99
		SEQ ID NO:153		SEQ ID NO:154				
miR-126	miR-126GSP	CATGATCAGCTGGGCCAAGAGCATTATTAC	miR-126RP	T+CG+TACCGTGAGTA	-0.26	8.937	0.18	1.80
		SEQ ID NO:155		SEQ ID NO:156				
miR-126*	miR-126*GSP3	CATGATCAGCTGGGCCAAGACGCGTACC	miR-126*RP	C+ATT+ATTA+CTTTTGGTA	-0.2969	8.184	3.58	35.78
		SEQ ID NO:157		SEQ ID NO:158				
miR-127	miR-127GSP	CATGATCAGCTGGGCCAAGAAGCCAAGCTC	miR-127RP	T+CG+GATCCGTCTGA	-0.2432	9.1013	1.11	11.13
		SEQ ID NO:159		SEQ ID NO:160				
miR-128a	miR-128aGSP	CATGATCAGCTGGGCCAAGAAAAAGAGACC	miR-128aRP	T+CA+CAGTGAACCGG	-0.2866	8.0867	0.16	1.60
		SEQ ID NO:161		SEQ ID NO:162				

IR-128b Extension frimer sequence iiR-128b miR-128bGSP CATGATCAGCTGGGCCAAGAGAAGAGCCAG miR-129 miR-129GSP CATGATCAGCTGGGCCAAGAGAGCCAGG miR-130a miR-130aGSP CATGATCAGCTGGGCCAAGAATGCCCTTT miR-130b miR-130bGSP CATGATCAGCTGGGCCAAGAATGCCCTTT miR-133a miR-133aGSP CATGATCAGCTGGGCCAAGAATGCCCTTT miR-133a miR-133aGSP CATGATCAGCTGGGCCAAGAATGCCCTTT miR-133a miR-133aGSP CATGATCAGCTGGGCCAAGAATGCCTTT sEQ ID NO:171 sEQ ID NO:173 miR-134 miR-134GSP CATGATCAGCTGGGCCAAGACAGCTGGTT sEQ ID NO:175 sEQ ID NO:177 miR-135a miR-135aGSP CATGATCAGCTGGGCCAAGACCTTGGTC sEQ ID NO:177 sEQ ID NO:177 miR-135bGSP CATGATCAGCTGGGCCAAGACCATAGGA sEQ ID NO:177 sEQ ID NO:177 miR-135bGSP CATGATCAGCTGGGCCAAGACACATAGGA sEQ ID NO:177 sEQ ID NO:181 miR-136GSP CATGATCAGCTGGGCCAAGACACACACACACACACACACA		5	Reverse Primer		5		Background RNA input	round
miR-129GSP miR-130aGSP miR-133aGSP miR-133aGSP miR-135aGSP miR-135aGSP miR-135aGSP		TGGGCCAAGAGAAAGAGCC	miR-128bRP	T+CA+CAGTGAACCGG	-0.2923	8.0608	0.07	0.74
miR-129GSP miR-130bGSP miR-133aGSP miR-133bGSP miR-133bGSP miR-135aGSP miR-135aGSP miR-135aGSP	SEQ ID NO:163			SEQ ID NO:164				
a miR-1306GSP miR-1336GSP miR-1336GSP miR-1356GSP miR-1356GSP miR-1356GSP miR-1356GSP		TGGGCCAAGAGCAAGCCCAG	miR-129RP	CTTTT+TG+CGGTCTG	-0.2942	9.7731	0.88	8.85
a miR-130aGSP miR-132GSP miR-133aGSP miR-133bGSP miR-135aGSP miR-135aGSP miR-135aGSP	SEQ ID NO:165			SEQ ID NO:166				
a miR-1356SP miR-1336GSP miR-1356GSP miR-1356GSP miR-1356GSP miR-1356GSP		TIGGGCCAAGAATGCCCTTTT	miR-130aRP	C+AG+TGCAATGTTAAAAG	-0.2943	8.7465	1.28	12.78
a miR-1356SP miR-1336GSP miR-1336GSP miR-1356GSP miR-1356GSP miR-1356GSP miR-1356GSP	SEQ ID NO:167			SEQ ID NO:168				
miR-132GSP miR-1336GSP miR-135GSP miR-1356GSP miR-1356GSP		TIGGGCCAAGAATGCCCTTTC	miR-130bRP	C+AG+TGCAATGATGA	-0.2377	9.1403	3.14	31.44
miR-132GSP miR-133aGSP miR-134GSP a miR-135aGSP miR-135aGSP miR-135bGSP	SEQ ID NO:169			SEQ ID NO:170				
a miR-133aGSP miR-134GSP miR-135aGSP miR-135bGSP miR-135bGSP		TIGGGCCAAGACGACCATGGC	miR-132RP	T+AA+CAGTCTACAGCC	-0.2948	8.1167	0.11	1.13
miR-133aGSP miR-134GSP miR-135aGSP miR-135bGSP miR-135bGSP	SEQ ID NO:171			SEQ ID NO:172				
miR-1356SP miR-1356SP miR-1356GSP miR-1356GSP		TTGGGCCAAGAACAGCTGGTT	miR-133aRP	T+TG+GTCCCCTTCAA	-0.295	9.3679	0.10	1.04
a miR-135bGSP a miR-135aGSP b miR-135bGSP miR-135bGSP	SEQ ID NO:173			SEQ ID NO:174				
miR-134GSP miR-135aGSP miR-135bGSP miR-136GSP		TGGGCCAAGATAGCTGGTTG	miR-133bRP	T+TG+GTCCCCTTCAA	-0.3062	8.3649	0.02	0.18
miR-134GSP miR-135aGSP miR-135bGSP miR-136GSP	SEQ ID NO:175			SEQ ID NO:176				
miR-135aGSP miR-135bGSP miR-136GSP		TGGGCCAAGACCCTCTGGTC	miR-134RP	T+GT+GACTGGTTGAC	-0.2965	9.0483	0.14	1.39
miR-135aGSP miR-135bGSP miR-136GSP	SEQ ID NO:177			SEQ ID NO:178				
b miR-135bGSP miR-136GSP		TGGGCCAAGATCACATAGGA	miR-135aRP	T+AT+GGCTTTTTATTCCT	-0.2914	8.092	1.75	17.50
b miR-135bGSP miR-136GSP	SEQ ID NO:179			SEQ ID NO:180				
miR-136GSP		TGGGCCAAGACACATAGGAA	miR-135bRP	T+AT+GGCTTTTCATTCC	-0.2962	7.8986	0.05	0.49
miR-136GSP	SEQ ID NO:181			SEQ ID NO:182				
SEQ ID NO:183		TGGGCCAAGATCCATCATCA	miR-136RP	A+CT+CCATTTGTTTTGATG	-0.3616	10.229	89.0	6.77
	SEQ ID NO:183			SEQ ID NO:184				
miR-137 miR-137GSP CATGATCAGCTGGGCCAAGACTACGCGTAT		CTGGGCCAAGACTACGCGTAT	miR-137RP	T+AT+TGCTTAAGAATACGC	-0.2876	8.234	8.57	85.71
SEQ ID NO:185	SEQ ID NO:185			SEQ ID NO:186				

Цитоп								
Target micro	Extension		Reverse				Backg	Background DNA input
RNA	Primer Name	Extension Primer Sequence	Name	Reverse Primer Sequence	Slope	Intercept	50ug	impur Sug
iR-138	miR-138GSP2	CATGATCAGCTGGGCCAAGACGGCCTGAT SEO ID NO:187	miR-138RP	A+GC+TGGTGTTGTGA SFO ID NO-188	-0.3023	9.0814	0.22	2.19
ıiR-139	miR-139GSP	CATGATCAGCTGGGCCAAGAAGACACGTGC	miR-139RP	T+CT+ACAGTGCACGT	-0.2983	8.1141	6.92	69.21
		SEQ ID NO:189		SEQ ID NO:190				
miR-140	miR-140GSP	CATGATCAGCTGGGCCAAGACTACCATAGG	miR-140RP	A+GT+GGTTTTACCCT	-0.2312	8.3231	0.13	1.34
		SEQ ID NO:191		SEQ ID NO:192				
miR-141	miR-141GSP9#	CATGATCAGCTGGGCCAAGACCATCTTTA	miR-	TAA+CAC+TGTCTGGTAA	-0.2805	9.6671	0.13	1.26
		SEQ ID NO:193	141RP2#	SEQ ID NO:194				
miR-142-	miR-142-	CATGATCAGCTGGGCCAAGATCCATAAA	miR-142-	TGT+AG+TGTTTCCTACT	-0.2976	8.4046	0.03	0.27
3p	3pGSP3	SEQ ID NO:195	3pRP	SEQ ID NO:196				
miR-143	miR-143GSP8#	CATGATCAGCTGGGCCAAGATGAGCTAC	miR-	T+GA+GATGAAGCACTG	-0.3008	9.2675	0.37	3.71
		SEQ ID NO:197	143RP2#	SEQ ID NO:198				
miR-144	miR-144GSP2	CATGATCAGCTGGGCCAAGACTAGTACAT	miR-144RP	TA+CA+GTAT+AGATGATG	-0.2407	9.4441	0.95	9.52
		SEQ ID NO:199		SEQ ID NO:200				
miR-145	miR-145GSP2	CATGATCAGCTGGGCCAAGAAAGGGATTC	miR-145RP	G+TC+CAGTTTTCCCA	-0.2937	8.0791	0.39	3.86
		SEQ ID NO:201		SEQ ID NO:202				
miR-146	miR-146GSP3	CATGATCAGCTGGGCCAAGAAACCCATG	miR-146RP	T+GA+GAACTGAATTCCA	-0.2861	8.8246	80.0	0.75
		SEQ ID NO:203		SEQ ID NO:204				
miR-147	miR-147GSP	CATGATCAGCTGGGCCAAGAGCAGAAGCAT	miR-147RP	G+TG+TGTGGAAATGC	-0.2989	8.8866	1.65	16.47
		SEQ ID NO:205		SEQ ID NO:206	•			
miR-148a	miR-148aGSP2	CATGATCAGCTGGGCCAAGAACAAAGTTC	miR-	T+CA+GTGCACTACAGAACT	-0.2928	9.4654	1.27	12.65
		SEQ ID NO:207	148aRP2	SEQ ID NO:208				
miR-148b	miR-148bGSP2	CATGATCAGCTGGGCCAAGAACAAAGTTC	miR-148bRP	T+CA+GTGCATCACAG	-0.2982	10.417	0.24	2.44
		SEQ ID NO:209		SEQ ID NO:210			÷	

Human								,
I arget micro RNA	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Slope	Intercept	Backgr RNA in 50ug	Background RNA input 50ug 5ug
iR-149	miR-149GSP2	CATGATCAGCTGGGCCAAGAGGAGTGAAG	miR-149RP	T+CT+GGCTCCGTGTC	-0.2996	8.3392	2.15	21.50
		SEQ ID NO:211		SEQ ID NO:212				
iR-150	miR-150GSP3	CATGATCAGCTGGGCCAAGACACTGGTA	miR-150RP	T+CT+CCCAACCCTTG	-0.2943	8.3945	90.0	0.56
		SEQ ID NO:213		SEQ ID NO:214				
miR-151	miR-151GSP2	CATGATCAGCTGGGCCAAGACCTCAAGGA	miR-151RP	A+CT+AGACTGAAGCTC	-0.2975	8.651	0.16	1.60
		SEQ ID NO:215		SEQ ID NO:216				
miR-152	miR-152GSP2	CATGATCAGCTGGGCCAAGACCCAAGTTC	miR-152RP	T+CA+GTGCATGACAG	-0.2741	8.7404	0.33	3.25
		SEQ ID NO:217		SEQ ID NO:218				
miR-153	miR-153GSP2	CATGATCAGCTGGGCCAAGATCACTTTTG	miR-153RP	TTG+CAT+AGTCACAAAA	-0.2723	9.5732	3.32	33.19
		SEQ ID NO:219		SEQ ID NO:220		٠		
miR-154*	miR-	CATGATCAGCTGGGCCAAGAAATAGGTCA	miR-	AATCA+TA+CACGGTTGAC	-0.3056	8.8502	0.07	0.74
	154*GSP9#	SEQ ID NO:221	154*RP2#	SEQ ID NO:222				
miR-154	miR-154GSP9#	CATGATCAGCTGGGCCAAGACGAAGGCAA	miR-	TA+GGTTA+TCCGTGTT	-0.3062	9.3947	0.10	96.0
		SEQ ID NO:223	154RP3#	SEQ ID NO:224				
miR-155	miR-155GSP8#	CATGATCAGCTGGGCCAAGACCCCTATC	miR-	TT+AA+TGCTAATCGTGATA	-0.3201	8.474	5.49	54.91
		SEQ ID NO:225	155RP2#	GG SEO ID NO:226				
miR-181a	miR-	CATGATCAGCTGGGCCAAGAACTCACCGA	miR-	AA+CATT+CAACGCTGTC	-0.2919	7.968	1.70	17.05
	181aGSP9#	SEQ ID NO:227	181aRP2#	SEQ ID NO:228				
miR-181c	miR-	CATGATCAGCTGGGCCAAGAACTCACCGA	miR-	AA+CATT+CAACCTGTCG	-0.3102	7.9029	1.08	10.78
	181cGSP9#	SEQ ID NO:229	181cRP2#	SEQ ID NO:230				
miR-182*	miR-182*GSP	CATGATCAGCTGGGCCAAGATAGTTGGCAA	miR-182*RP	T+GG+TTCTAGACTTGC	-0.2978	8.5876	4.25	42.47
		SEQ ID NO:231		SEQ ID NO:232				
miR-182	miR-182GSP2	CATGATCAGCTGGGCCAAGATGTGAGTTC	miR-182RP	TTT+GG+CAATGGTAG	-0.2863	9.0854	1.52	15.20
		SEQ ID NO:233		SEQ ID NO:234				

Human Target micro RNA	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Slope	Intercept	Backgr RNA i	Background RNA input 50ug 5ug
iR-183	miR-183GSP2	CATGATCAGCTGGGCCAAGACAGTGAATT SEQ ID NO:235	miR-183RP	T+AT+GGCACTGGTAG SEQ ID NO:236	-0.2774	9.9254	1.95	19.51
iR-184	miR-184GSP2	CATGATCAGCTGGGCCAAGAACCCTTATC SEQ ID NO:237	miR-184RP	T+GG+ACGGAGAACTG SEQ ID NO:238	-0.2906	7.9585	0.05	0.49
miR-186	miR-186GSP9#	CATGATCAGCTGGGCCAAGAAAGCCCAAA SEQ ID NO:239	miR- 186RP3#	CA+AA+GAATT+CTCCTTTT GG SEQ ID NO:240	-0.2861	8.6152	0.32	3.18
miR-187	miR-187GSP	CATGATCAGCTGGGCCAAGACGGCTGCAAC SEQ ID NO:241	miR-187RP	T+CG+TGTCTTGTGTT SEQ ID NO:242	-0.2953	7.9329	1.23	12.31
miR-188	miR-188GSP	CATGATCAGCTGGGCCAAGAACCCTCCACC SEQ ID NO:243	miR-188RP	C+AT+CCCTTGCATGG SEQ ID NO:244	-0.2925	8.0782	8.49	84.92
miR-189	miR-189GSP2	CATGATCAGCTGGGCCAAGAACTGATATC SEQ ID NO:245	miR-189RP	G+TG+CCTACTGAGCT SEQ ID NO:246	-0.2981	8.8964	0.21	2.08
miR-190	miR-190GSP9#	CATGATCAGCTGGGCCAAGAACCTAATAT SEQ ID NO:247	miR- 190RP4#	T+GA+TA+TGTTTGATATAT TAG SEQ ID NO:248	-0.3317	9.8766	0.43	4.34
miR-191	miR-191GSP2	CATGATCAGCTGGGCCAAGAAGCTGCTTT SEQ ID NO:249	miR-191RP2	C+AA+CGGAATCCCAAAAG SEQ ID NO:250	-0.299	9.0317	0.41	4.07
miR-192	miR-192GSP2	CATGATCAGCTGGGCCAAGAGGCTGTCAA SEQ ID NO:251	miR-192RP	C+TGA+CCTATGAATTGAC SEQ ID NO:252	-0.2924	9.5012	1.10	10.98
miR-193	miR-193GSP9#	CATGATCAGCTGGGCCAAGACTGGGACTT SEQ ID NO:253	miR- 193RP2#	AA+CT+GGCCTACAAAG SEQ ID NO:254	-0.3183	8.9942	0.17	1.72
miR-194	mir194GSP8#	CATGATCAGCTGGGCCAAGATCCACATG SEQ ID NO:255	mir194RP#	TG+TAA+CAGCAACTCCA SEQ ID NO:256	-0.3078	8.8045	0.37	3.69

KINA ip 105	Extension	, ,	Reverse Primer		ā	, , , , , , , , , , , , , , , , , , ,	Background RNA input	round input
	miR-195GSP9#	CATGATCAGCTGGGCCAAGACCAATATT	miR-	T+AG+CAG+CACAGAAATA	-0.2955	10.213	97.0	7.58
		SEQ ID NO:257	195RP3#	SEQ ID NO:258				
IR-196b	miR-196bGSP	CATGATCAGCTGGGCCAAGACCAACAG	miR-196bRP	TA+GGT+AGTTTCCTGT	-0.301	8.1641	1.47	14.66
		SEQ ID NO:259		SEQ ID NO:260				
miR-196a 1	miR-196aGSP	CATGATCAGCTGGGCCAAGACCAACAT	miR-196aRP	TA+GG+TAGTTTCATGTTG	-0.2932	8.0448	8.04	80.37
		SEQ ID NO:261		SEQ ID NO:262				
miR-197 I	miR-197GSP2	CATGATCAGCTGGGCCAAGAGCTGGGTGG	miR-197RP	TT+CA+CCACCTTCTC	-0.289	8.2822	0.71	7.10
		SEQ ID NO:263		SEQ ID NO:264				
miR-198 I	miR-198GSP3	CATGATCAGCTGGGCCAAGACCTATCTC	miR-198RP	G+GT+CCAGAGGGAG	-0.2986	8.1359	0.31	3.15
		SEQ ID NO:265		SEQ ID NO:266				
	miR-	CATGATCAGCTGGGCCAAGAAACCAATGT	miR-	T+AC+AGTAGTCTGCAC	-0.3029	9.0509	0.25	2.52
199a*]	199a*GSP2	SEQ ID NO:267	199a*RP	SEQ ID NO:268				
miR-199a 1	miR-199aGSP2	CATGATCAGCTGGGCCAAGAGAACAGGTA	miR-199aRP	C+CC+AGTGTTCAGAC	-0.3187	9.2268	0.12	1.16
		SEQ ID NO:269		SEQ ID NO:270				
miR-199b 1	miR-199bGSP	CATGATCAGCTGGGCCAAGAGAACAGATAG	miR-199bRP	C+CC+AGTGTTTAGAC	-0.3165	9.3935	2.00	20.04
		SEQ ID NO:271		SEQ ID NO:2/2				
miR-200a	miR-200aGSP2	CATGATCAGCTGGGCCAAGAACATCGTTA	miR-200aRP	TAA+CAC+TGTCTGGT	-0.2754	9.1227	80.0	0.78
		SEQ ID NO:273		SEQ ID NO:274				_
miR-200b	miR-200bGSP2	CATGATCAGCTGGGCCAAGAGTCATCATT	miR-200bRP	TAATA+CTG+CCTGGTAAT	-0.2935	8.5461	80.0	0.85
		SEQ ID NO:275		SEQ ID NO:276				
miR-202	miR-202	CATGATCAGCTGGGCCAAGATTTTCCCATG	miR-202RP#	A+GA+GGTATA+GGGCAT	-0.2684	9:026	0.25	2.48
	GSP10#	SEQ ID NO:277		SEQ ID NO:278				
miR-203 1	miR-203GSP2	CATGATCAGCTGGGCCAAGACTAGTGGTC	miR-203RP	G+TG+AAATGTTTAGGACC	-0.2852	8.1279	1.60	16.03
		SEQ ID NO:279		SEQ ID NO:280				

Human			Ravorsa				Barko	Background
micro RNA	Extension Primer Name	Extension Primer Sequence	Primer Name	Reverse Primer Sequence	Slope	Intercept	RNA 50ug	RNA input 50ug 5ug
iiR-204	miR-204GSP2	CATGATCAGCTGGGCCAAGAAGGCATAGG SEQ ID NO:281	miR-204RP	T+TC+CCTTTGTCATCC SEQ ID NO:282	-0.2925	8.7648	0.16	1.59
iR-205	miR-205GSP	CATGATCAGCTGGGCCAAGACAGACTCCGG SEQ ID NO:283	miR-205RP	T+CCTT+CATTCCACC SEQ ID NO:284	-0.304	8.2407	9.21	92.15
miR-206	mir206GSP7#	CATGATCAGCTGGGCCAAGACCACACA SEQ ID NO:285	miR-206RP#	T+G+GAA+TGTAAGGAAGT GT SEQ ID NO:286	-0.2815	8.2206	0.29	2.86
miR-208	miR- 208_GSP13#	CATGATCAGCTGGGCCAAGAACAAGCTTTTTGC SEQ ID NO:287	miR- 208_RP4#	ATAA+GA+CG+AGCAAAAA G SEQ ID NO:288	-0.2072	7.9097	57.75	577.52
miR-210	miR-210GSP	CATGATCAGCTGGGCCAAGATCAGCCGCTG SEQ ID NO:289	miR-210RP	C+TG+TGCGTGTGACA SEQ ID NO:290	-0.2717	8.249	0.18	1.77
miR-211	miR-211GSP2	CATGATCAGCTGGGCCAAGAAGGCGAAGG SEQ ID NO:291	miR-211RP	T+TC+CCTTTGTCATCC SEQ ID NO:292	-0.2926	8.3106	0.10	1.00
miR-212	miR-212GSP9#	CATGATCAGCTGGGCCAAGAGGCCGTGAC SEQ ID NO:293	miR- 212RP2#	T+AA+CAGTCTCCAGTCA SEQ ID NO:294	-0.2916	8.0745	0.59	5.86
miR-213	miR-213GSP	CATGATCAGCTGGGCCAAGAGGTACAATCA SEQ ID NO:295	miR-213RP	A+CC+ATCGACCGTTG SEQ ID NO:296	-0.2934	8.1848	2.96	29.59
miR-214	miR-214GSP	CATGATCAGCTGGGCCAAGACTGCCTGTCT SEQ ID NO:297	miR-214RP	A+CA+GCAGGCACAGA SEQ ID NO:298	-0.2947	7.82	0.84	8.44
miR-215	miR-215GSP2	CATGATCAGCTGGGCCAAGAGTCTGTCAA SEQ ID NO:299	miR-215RP	A+TGA+CCTATGAATTGAC SEQ ID NO:300	-0.2932	8.9273	1.51	15.05
miR-216	miR-216GSP9#	CATGATCAGCTGGGCCAAGACACAGTTGC SEQ ID NO:301	mir216RP#	TAA+TCT+CAGCTGGCA SEQ ID NO:302	-0.273	8.5829	0.95	9.50

micro RNA	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Slope	Intercept	Backgr RNA is 50ug	Background RNA input 50ug 5ug
iiR-217	miR-217GSP2	CATGATCAGCTGGGCCAAGAATCCAATCA	miR-217RP2	T+AC+TGCATCAGGAACTGA	-0.3089	9.6502	0.07	0.71
		SEQ ID NO:303		SEQ ID NO:304				
iIR-218	miR-218GSP2	CATGATCAGCTGGGCCAAGAACATGGTTA	miR-218RP	TTG+TGCTT+GATCTAAC	-0.2778	8.4363	1.00	10.05
		SEQ ID NO:305		SEQ ID NO:306		-		
miR-220	miR-220GSP	CATGATCAGCTGGGCCAAGAAAAGTGTCAG	miR-220RP	C+CA+CACCGTATCTG	-0.2755	9.0728	8.88	88.75
		SEQ ID NO:307		SEQ ID NO:308				
miR-221	miR-221GSP9#	CATGATCAGCTGGGCCAAGAGAAACCCAG	miR-221RP#	A+GC+TACATTGTCTGC	-0.2886	8.5743	0.12	1.17
		SEQ ID NO:309		SEQ ID NO:310				
miR-222	miR-222GSP8#	CATGATCAGCTGGGCCAAGAGAGACCCA	miR-222RP#	A+GC+TACATCTGGCT	-0.283	8.91	1.64	16.41
		SEQ ID NO:311		SEQ ID NO:312				
miR-223	miR-223GSP	CATGATCAGCTGGGCCAAGAGGGGTATTTG	miR-223RP	TG+TC+AGTTTGTCAAA	-0.2998	8.6669	0.94	9.44
		SEQ ID NO:313		SEQ ID NO:314				
miR-224	miR-224GSP8#	CATGATCAGCTGGGCCAAGATAAACGGA	miR-	C+AAG+TCACTAGTGGTT	-0.2802	7.5575	0.56	5.63
		SEQ ID NO:315	224RP2#	SEQ ID NO:316				
miR-296	miR-296GSP9#	CATGATCAGCTGGGCCAAGAACAGGATTG	miR-	A+GG+GCCCCCCTCAA	-0.3178	8.3856	0.10	96.0
		SEQ ID NO:317	296RP2#	SEQ ID NO:318				
miR-299	miR-299GSP9#	CATGATCAGCTGGGCCAAGAATGTATGTG	miR-299RP#	T+GG+TTTACCGTCCC	-0.3155	7.9383	1.30	12.96
		SEQ ID NO:319		SEQ ID NO:320				
miR-301	miR-301GSP	CATGATCAGCTGGGCCAAGAGCTTTGACAA	miR-301RP	C+AG+TGCAATAGTATTGT	-0.2839	8.314	2.55	25.52
		SEQ ID NO:321		SEQ ID NO:322		:		
miR-	miR-302a*GSP	CATGATCAGCTGGGCCAAGAAAAGCAAGTA	miR-	TAAA+CG+TGGATGTAC	-0.2608	8.3921	0.04	0.41
302a*		SEQ ID NO:323	302a*RP	SEQ ID NO:324				
miR-302a	miR-302aGSP	CATGATCAGCTGGGCCAAGATCACCAAAAC	miR-302aRP	T+AAG+TGCTTCCATGT	-0.2577	9.6657	2.17	21.67
		SEQ ID NO:325		SEQ ID NO:326				

Human Target micro	Extension		Reverse				Backg	Background RNA innut
RNA	Primer Name	Extension Primer Sequence	Name	Reverse Primer Sequence	Slope	Intercept	50ug	50ug Sug
iiR- 02b*	miR-302b*GSP	CATGATCAGCTGGGCCAAGAAGAAGCACT SEQ ID N0:327	miR- 302b*RP	A+CTTTAA+CATGGAAGTG SEQ ID NO:328	-0.2702	8.5153	0.02	0.24
iR-302b	miR-302bGSP	CATGATCAGCTGGGCCAAGACTACTAAAAC SEO ID NO:329	miR-302bRP	T+AAG+TGCTTCCATGT SEO ID NO:330	-0.2398	9.1459	5.11	51.11
miR-302d	miR-302dGSP	CATGATCAGCTGGGCCAAGAACACTCAAAC SEQ ID NO:331	miR-302dRP	T+AAG+TGCTTCCATGT SEQ ID NO:332	-0.2368	8.5602	5.98	59.78
miR- 302c*	miR- 302c*_GSP9#	CATGATCAGCTGGGCCAAGACAGCAGGTA SEQ ID NO:333	miR- 302c*_RP2#	TT+TAA+CAT+GGGGGTACC SEQ ID NO:334	-0.312	8.2904	0.33	3.28
miR-302c	miR- 302cGSP9#	CATGATCAGCTGGGCCAAGACCACTGAAA SEQ ID NO:335	miR- 302cRP5#	T+AAG+TGCTTCCATGTTTC A SEQ ID NO:336	-0.2945	8.381	14.28	142.76
miR-320	miR- 320_GSP8#	CATGATCAGCTGGGCCAAGATTCGCCCT SEQ ID NO:337	miR- 320_RP3#	AAAA+GCT+GGGTTGAGAG G SEQ ID NO:338	-0.2677	7.8956	6.73	67.29
miR-323	miR-323GSP	CATGATCAGCTGGGCCAAGAAGAGGTCGAC SEQ ID NO:339	miR-323RP	G+CA+CATTACACGGT SEQ ID NO:340	-0.2878	8.2546	0.19	1.92
miR-324- 3p	miR-324- 3pGSP	CATGATCAGCTGGGCCAAGACCAGCAGCAC SEQ ID NO:341	miR-324- 3pRP	C+CA+CTGCCCCAGGT SEQ ID NO:342	-0.2698	8.5223	2.54	25.41
miR-324- 5p	miR-324- 5pGSP	CATGATCAGCTGGGCCAAGAACACCAATGC SEQ ID NO:343	miR-324- 5pRP	C+GC+ATCCCCTAGGG SEQ ID NO:344	-0.2861	7.6865	90.00	0.62
miR-325	miR-325GSP	CATGATCAGCTGGGCCAAGAACACTTACTG SEQ ID NO:345	miR-325RP	C+CT+AGTAGGTGTCC SEQ ID NO:346	-0.2976	8.1925	0.01	0.14
miR-326	miR-326GSP	CATGATCAGCTGGGCCAAGACTGGAGGAAG SEQ ID N0:347	miR-326RP	C+CT+CTGGGCCCTTC SEQ ID NO:348	-0.2806	7.897	0.59	5.87

Human Target	7-7-7-1		Reverse				Backg	Background
RNA	Extension Primer Name	Extension Primer Sequence	Primer Name	Reverse Primer Sequence	Slope	Intercept	50ug Sug	input Sug
iiR-328	miR-328GSP	CATGATCAGCTGGGCCAAGAACGGAAGGGC SEQ ID NO:349	miR-328RP	C+TG+GCCCTCTCTGC SEQ ID NO:350	-0.293	7.929	3.17	31.69
uiR-330	miR-330GSP	CATGATCAGCTGGGCCAAGATCTCTGCAGG SEQ ID NO:351	miR-330RP	G+CA+AAGCACACGGC SEQ ID NO:352	-0.3009	7.7999	0.13	1.30
miR-331	miR-331GSP	CATGATCAGCTGGGCCAAGATTCTAGGATA SEQ ID NO:353	miR-331RP	G+CC+CCTGGGCCTAT SEQ ID NO:354	-0.2816	8.1643	0.45	4.54
miR-337	miR-337GSP	CATGATCAGCTGGGCCAAGAAAGGCATCA SEQ ID NO:355	miR-337RP	T+CC+AGCTCCTATATG SEQ ID NO:356	-0.2968	8.7313	0.10	1.02
miR-338	miR-338GSP	CATGATCAGCTGGGCCAAGATCAACAAAAT SEQ ID NO:357	miR-338RP2	T+CC+AGCATCAGTGATTT SEQ ID NO:358	-0.2768	8.5618	0.52	5.17
miR-339	miR-339GSP9#	CATGATCAGCTGGGCCAAGATGAGCTCCT SEQ ID NO:359	miR- 339RP2#	T+CC+CTGTCCTCCAGG SEQ ID NO:360	-0.303	8.4873	0.27	2.72
miR-340	miR-340GSP	CATGATCAGCTGGGCCAAGAGGCTATAAAG SEQ ID NO:361	miR-340RP	TC+CG+TCTCAGTTAC SEQ ID NO:362	-0.2846	9.6673	0.15	1.45
miR-342	miR-342GSP3	CATGATCAGCTGGGCCAAGAGACGGGTG SEQ ID NO:363	miR-342RP	T+CT+CACACAGAAATCG SEQ ID NO:364	-0.293	8.1553	4.69	46.85
miR-345	miR-345GSP	CATGATCAGCTGGGCCAAGAGCCCTGGACT SEQ ID NO:365	miR-345RP	T+GC+TGACTCCTAGT SEQ ID NO:366	-0.2909	8.468	0.04	0.40
miR-346	miR-346GSP	CATGATCAGCTGGGCCAAGAAGAGGCAGGC SEQ ID NO:367	miR-346RP	T+GT+CTGCCCGCATG SEQ ID NO:368	-0.2959	8.1958	0.25	2.54
miR-363	miR-363 GSP10#	CATGATCAGCTGGGCCAAGATACAGATGGA SEQ ID NO:369	miR-363RP#	AAT+TG+CAC+GGTATCC SEQ ID NO:370	-0.2362	8.9762	0.44	4.36
miR-367	miR-367GSP	CATGATCAGCTGGGCCAAGATCACCATTGC SEQ ID NO:371	miR-367RP	AAT+TG+CACTTTAGCAAT SEQ ID NO:372	-0.2819	8.6711	0.00	0.03

Human			ŕ				,	,
l arget micro	Extension		Reverse Primer				Backg RNA	Background RNA input
RNA	Primer Name	Extension Primer Sequence	Name	Reverse Primer Sequence	Slope	Intercept	20ug	5ug
iiR-368	miR-368GSP	CATGATCAGCTGGGCCAAGAAAACGTGGAA SEO ID NO:373	miR-368RP2	A+CATAGA+GGAAATTCCA C	-0.2953	8.0067	6.01	60.11
				SEQ ID NO:374				
miR-370	miR-370GSP	CATGATCAGCTGGGCCAAGACCAGGTTCCA	miR-370RP	G+CC+TGCTGGGGTGG	-0.2825	8.3162	1.45	14.55
		SEQ ID NO:375		SEQ ID NO:376				
miR-371	miR-371GSP	CATGATCAGCTGGGCCAAGAACACTCAAAA	miR-371RP	G+TG+CCGCCATCTTT	-0.295	7.8812	2.51	25.12
		SEQ ID NO:377		SEQ ID NO:378				
miR-372	miR-372GSP	CATGATCAGCTGGGCCAAGAACGCTCAAAT	miR-372RP	A+AA+GTGCTGCGACA	-0.2984	8.9183	0.05	0.53
		SEQ ID NO:379		SEQ ID NO:380				
miR-373*	miR-373*GSP	CATGATCAGCTGGGCCAAGAGGAAAGCGCC	miR-373*RP	A+CT+CAAAATGGGGG	-0.2705	8.4513	0.20	1.99
		SEQ ID NO:381		SEQ ID NO:382				
miR-373	miR-373GSP	CATGATCAGCTGGGCCAAGAACACCCCAAA	miR-373RP2	GA+AG+TGCTTCGATTTTGG	-0.307	7.9056	9.13	91.32
		SEQ ID NO:383		SEQ ID NO:384				
miR-374	miR-374GSP2	CATGATCAGCTGGGCCAAGACACTTATCA	miR-374RP	TT+AT+AATA+CAACCTGAT	-0.2655	9.3795	9.16	91.60
		SEQ ID NO:385		AAG				
				SEQ ID NO:386				
miR-375	miR-375GSP	CATGATCAGCTGGGCCAAGATCACGCGAGC	miR-375RP	TI+TG+TTCGTTCGGC	-0.3041	8.1181	0.09	0.90
		SEQ ID NO:387		SEQ ID NO:388				
miR-376b	miR-376b	CATGATCAGCTGGGCCAAGAAACATGGA	miR-	AT+CAT+AGA+GGAAAATCC	-0.2934	9.0188	1.07	10.74
	GSP8#	SEQ ID NO:389	376bRP#	A				
				SEQ ID NO:390				
miR-378	miR-378GSP	CATGATCAGCTGGGCCAAGAACACAGGACC	miR-378RP	C+TC+CTGACTCCAGG	-0.2899	8.1467	0.07	0.73
		SEQ ID NO:391		SEQ ID NO:392				
miR-379	miR-	CATGATCAGCTGGGCCAAGATACGTTC	miR-	T+GGT+AGACTATGGAACG	-0.2902	8.2149	10.89	108.86
	379_GSP7#	SEQ ID NO:393	379RP2#	SEQ ID NO:394				

Juman Target	7		Reverse				Background RNA input	ound
RNA	Extension Primer Name	Extension Primer Sequence	Name	Reverse Primer Sequence	Slope	Intercept	50ug	5ug
iR-380-	miR-380- 5pGSP	CATGATCAGCTGGGCCAAGAGCGCATGTTC SEQ ID NO:395	miR-380- 5pRP	T+GGT+TGACCATAGA SEQ ID NO:396	-0.2462	9.4324	1.30	13.04
iR-380- 3p	miR-380- 3pGSP	CATGATCAGCTGGGCCAAGAAAGATGTGGA SEQ ID NO:397	miR-380- 3pRP	TA+TG+TAATATGTCCACA SEQ ID NO:398	-0.3037	8.0356	3.69	36.89
miR-381	miR-381GSP2	CATGATCAGCTGGGCCAAGAACAGAGAGC SEQ ID NO:399	miR-381RP2	TATA+CAA+GGGCAAGCT SEQ ID NO:400	-0.3064	8.8704	1.72	17.16
· miR-382	miR-382GSP	CATGATCAGCTGGGCCAAGACGAATCCACC SEQ ID NO:401	miR-382RP	G+AA+GTTGTTCGTGGT SEQ ID NO:402	-0.2803	7.6738	99.0	6.57
miR-383	miR-383GSP	CATGATCAGCTGGGCCAAGAAGCCACAATC SEQ ID NO:403	miR-383RP2	A+GATC+AGAAGGTGATTG T SEQ ID NO:404	-0.2866	8.1463	0.54	5.45
miR-410	miR-410 GSP9#	CATGATCAGCTGGGCCAAGAACAGGCCAT SEQ ID NO:405	miR-410RP#	AA+TA+TAA+CA+CAGATGG C SEQ ID NO:406	-0.2297	8.5166	4.27	42.71
miR-412	miR-412 GSP10#	CATGATCAGCTGGGCCAAGAACGGCTAGTG SEQ ID NO:407	miR-412RP#	A+CTT+CACCTGGTCCACTA SEQ ID NO:408	-0.3001	7.9099	4.24	42.37
miR-422a	miR-422aGSP	CATGATCAGCTGGGCCAAGAGGCCTTCTGA SEQ ID NO:409	miR-422aRP	C+TG+GACTTAGGGTC SEQ ID NO:410	-0.3079	9.3108	5.95	59.54
miR-422b	miR-422bGSP	CATGATCAGCTGGGCCAAGAGGCCTTCTGA SEQ ID NO:411	miR-4226RP	C+TG+GACTTGGAGTC SEQ ID NO:412	-0.2993	8.9437	4.86	48.56
miR-423	miR-423GSP	CATGATCAGCTGGGCCAAGACTGAGGGGCC SEQ ID NO:413	miR-423RP	A+GC+TCGGTCTGAGG SEQ ID NO:414	-0.3408	9.2274	90.9	60.62
miR-424	miR-424GSP#	CATGATCAGCTGGGCCAAGATTCAAAACAT SEQ ID NO:415	miR- 424RP2#	C+AG+CAGCAATTCATGTTT T SEQ ID NO:416	-0.3569	9.3419	10.78	107.85

747

icro Fri NA Pri -425 miR			Nevel Se				RNA input	nout
	Extension	Extension Primer Sequence	Primer Name	Reverse Primer Sequence	Slope	Intercept	50ug Sug	5ug
	miR-425GSP	TGGGCCAAG	miR-425RP	A+TC+GGGAATGTCGT	-0.2932	7.9786	0.39	3.93
		SEQ ID NO:417		SEQ ID NO:410		1000	,,,,	160 10
miR-429 miR-	- -	CATGATCAGCTGGGCCAAGAACGGTTTTACC	miR-	T+AATAC+TG+TCTGGTAAA A	-0.2458	8.2805	16.21	102.12
429	429_GSP11#	SEQ ID NO:419	429KL3"	SEQ ID NO:420				
miR-431 miF	miR-431	CATGATCAGCTGGGCCAAGATGCATGACGG	miR-431RP#	T+GT+CTTGCAGGCCG	-0.3107	7.7127	7.00	70.05
	GSP10#	SEQ ID NO:421		SEQ ID NO:422				
miR-448 miF	miR-448GSP	CATGATCAGCTGGGCCAAGAATGGGACATC	miR-448RP	TTG+CATA+TGTAGGATG	-0.3001	8.4969	0.12	1.16
		SEO ID NO:423		SEQ ID NO:424				
mip 440 miR-	-a	CATGATCAGCTGGGCCAAGAACCAGCTAAC	miR-	T+GG+CAGTGTATTGTTAGC	-0.3225	8.4953	2.57	25.70
	449GSP10#	SEO ID NO:425	449RP2#	SEQ ID NO:426				
miR-450 mil	miR-450GSP	CATGATCAGCTGGGCCAAGATATTAGGAAC	miR-450RP	TTTT+TG+CGATGTGTT	-0.2906	8.1404	0.48	4.82
		SEO ID NO.427		SEQ ID NO:428				
1		STATE OF A A A A A A A A A COLUMN A A A A A A A A A A A A A A A A A A A	#44.57	AAA+CCG+TTA+CCATTACT	-0.2544	8.0291	1.73	17.35
miR-451 mi	miR-451	CATGATCAGCTGGGCCAAGAAAACICAGTA	miK-451KF"	GA				
<u>ਤ</u>	GSF10"	SEQ ID NO:429		SEQ ID NO:430				
let7a 12+	12+70 GCD7#	CATGATCAGCTGGGCCAAGAAACTATAC	let7a-RP#	T+GA+GGTAGTAGGTTG	-0.3089	9.458	0.04	0.38
_	7 TG-071	SEQ ID NO:431		SEQ ID NO:432				
let7h	#60050 42+01	CATGATCAGCTGGGCCAAGAAACCACAC	let7b-RP#	T+GA+GGTAGTAGGTTG	-0.2978	7.9144	0.05	0.54
	7 100-0/1	SEO ID NO:433		SEQ ID NO:432				
let7c lat	#caso 25to1	CATGATCAGCTGGGCCAAGAAACCATAC	let7c-RP#	T+GA+GGTAGTAGGTTG	-0.308	7.9854	0.01	0.14
	7 100-27	SEO ID NO:434		SEQ ID NO:432				1
let7d	#0000 50	CATGATCAGCTGGGCCAAGAACTATGCA	let7d-RP#	A+GA+GGTAGTAGGTTG	-0.3238	8.3359	90.0	0.57
	161/a-GSF2"	SEO ID NO:435		SEQ ID NO:436				

Human			£				D. 2.1.2.0	
narget micro RNA	Extension Primer Name	Extension Primer Sequence	Keverse Primer Name	Reverse Primer Sequence	Slope	Intercept	Background RNA input 50ug 5ug	round Input Sug
:t7e	let7e-GSP2#	CATGATCAGCTGGGCCAAGAACTATACA SEQ ID NO:437	let7e-RP#	T+GA+GGTAGGAGGTTG SEQ ID NO:438	-0.3284	9.7594	0.22	2.20
JT.	let7f-GSP2#	CATGATCAGCTGGGCCAAGAAACTATAC SEQ ID NO:439	let7f-RP#	T+GA+GGTAGTAGATTG SEQ ID NO:440	-0.2901	11.107	0.32	3.18
let7g	let7g-GSP2#	CATGATCAGCTGGGCCAAGAACTGTACA SEQ ID NO:441	let7g-RP#	T+GA+GGTAGTAGTTTG SEQ ID NO:442	-0.3469	9.8235	0.16	1.64
let7i	let7i-GSP2#	CATGATCAGCTGGGCCAAGAACAGCACA SEQ ID NO:443	let7i-RP#	T+GA+GGTAGTAGTTTG SEQ ID NO:444	-0.321	10.82	0.20	1.99
miR-377	miR-377GSP	CATGATCAGCTGGGCCAAGAACAAAAGTTG SEQ ID NO:445	miR-377RP2	AT+CA+CACAAAGGCAAC SEQ ID NO:446	-0.2979	10.612	13.45	134.48
miR-376a	miR- 376a_GSP7	CATGATCAGCTGGGCCAAGAACGTGGA SEQ ID NO:447	miR- 376a_RP5	AT+CAT+AGA+GGAAAATCC SEQ ID NO:448	-0.2938	10.045	63.00	630.00
miR-22	miR-22GSP	CATGATCAGCTGGGCCAAGAACAGTTCTTC SEQ ID NO:449	miR-22RP	A+AG+CTGCCAGTTGA SEQ ID NO:450	-0.2862	8.883	20.46	204.58
miR-200c	miR-200cGSP2	CATGATCAGCTGGGCCAAGACCATCATTA SEQ ID NO:451	miR-200cRP	T+AA+TACTGCCGGGT SEQ ID NO:452	-0.3094	11.5	15.99	159.91
miR-24	miR-24GSP	CATGATCAGCTGGGCCAAGACTGTTCCTGC SEQ ID NO:453	miR-24RP	T+GG+CTCAGTTCAGC SEQ ID NO:454	-0.3123	8.6824	24.34	243.38
miR- 29cDNA	miR-29cGSP10	CATGATCAGCTGGGCCAAGAACCGATTTCA SEQ ID NO:455	miR-29cRP	T+AG+CACCATTTGAAAT SEQ ID NO:456	-0.2975	8.8441	23.22	232.17
miR-18	miR-18GSP	CATGATCAGCTGGGCCAAGATATCTGCACT SEQ ID NO:457	miR-18RP	T+AA+GGTGCATCTAGT SEQ ID NO:458	-0.3209	6660.6	14.90	149.01
miR-185	miR-185GSP	CATGATCAGCTGGGCCAAGAGAACTGCCTT SEQ ID NO:459	miR-185RP	T+GG+AGAGAAAGGCA SEQ ID NO:460	-0.3081	8.9289	15.73	157.32

								-
nan			Reverse				Background PNA innut	
jeg.	Extension		Primer	Reverse Primer Sequence	Slope	Intercept	50ug Sug	Sug
017	Drimor Nome	Extension Primer Sequence	Name	Metalso Firmer 2-4	0.011	10.046	15.87	158.67
1	rrimer traine	CATTACTTGGGCCAAGACCCACCGA	miR-	AA+CATT+CATTGCTGTC	0.3115	10.040		
181b	muk-	CALUATION OF THE CASE OF THE C	181bRP2#	SEQ ID NO:462				
-	1810GSF0	SEC ID INO:401		ECCO	AUMUO	annrox	approx.	approx.
_	100,000	CATGATCAGCTGGCCAAGAAAAAGAGACC	miR-	TCACAGTGAACCGG1	approx.	approus		1 60
miK-128a	mik-170a03r	Chiches	128anLRP	SEQ ID NO: 494	-0.2866	8.0807	0.10	
		SEQ ID NO:161	i	ACCTGGTGTTGTGAA	approx.	approx.	approx.	approx.
miR-138	miR-138GSP2	CATGATCAGCTGGGCCAAGACGGCCTGAT	mik- 138nl.RP	AUCTUATOR OF	-0.3023	9.0814	0.22	2.19
		SEO ID NO-187		SEQ ID NO. 455			Admino	annrox
		DATON OT A CONTROL OF THE PARTY	miR-	TGAGATGAAGCACTGT	approx.	approx.	approx.	approx.
miR-143	miR-143GSP8#	CATGATCAGCIGGCCAAGAIGACTAC	143nLRP	SEO ID NO: 496	-0.3008	9.2675	0.37	3.71
		SEO ID NO:197		מור אוני אוני) io	annrox	approx.
		ATOTOTO A CALCOCOCCOCACA CALCOCOCACA CALCOCACA	miR-	TCTCCCAACCCTTGTA	approx.	approx.	approv.	J. J. J. J.
miR-150	miR-150GSP3	CATGATCAGCIGGCCAAGACACIGGIA	150nLRP	SEO ID NO: 497	-0.2943	8.3945	90.0	0.56
		SEQ ID NO:213			annrox	approx.	approx.	approx.
3	7	CATGATCAGCTGGGCCAAGAACTCACCGA	miR-	AACAIICAACGCIGI	- Arker	277	1 70	17.05
miR-181a		CALCALOSTOS	181anLRP	SEO ID NO: 498	-0.2919	7.968	1./0	20.71
	181aGSP9#	SEQ ID NO:227		* OCCUPATION	onnrov	annrox.	approx.	approx.
		PLA JOHO COLOR A CATOLANA	miR-	TGTAACAGCAACICCA	approv.	and valde		
miR-194	mir194GSP8#	CATGATCAGCTGGGCCAACATTCGTTGGTTGGTTGGTTGG	194nLRP	SEO ID NO: 499	-0.3078	8.8045	0.37	3.09
		SEO ID NO:255						

EXAMPLE 4

This Example describes assays and primers designed for quantitative analysis of murine miNRA expression patterns.

Methods: The representative murine microRNA target templates described in TABLE 7 are publically available accessible on the World Wide Web at the Wellcome Trust Sanger Institute website in the "miRBase sequence database" as described in Griffith-Jones et al. (2004), Nucleic Acids Research 32:D109-D111 and and Griffith-Jones et al. (2006), Nucleic Acids Research 34: D140-D144. As indicated below in TABLE 7, the murine microRNA templates are either totally identical to the corresponding human microRNA templates, identical in the overlapping sequence with differing ends, or contain one or more base pair changes as compared to the human microRNA sequence. The murine microRNA templates that are identical or that have identical overlapping sequence to the corresponding human templates can be assayed using the same primer sets designed for the human microRNA templates, as indicated in TABLE 7. For the murine microRNA templates with one or more base pair changes in comparison to the corresponding human templates, primer sets have been designed specifically for detection of the murine microRNA, and these primers are provided in TABLE 7. The extension primer reaction and quantitative PCR reactions for detection of the murine microRNA templates may be carried out as described in EXAMPLE 3.

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TABLE 7: Primers to detect murine microRNA target templates

Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Mouse microRNA as compared to Human microRNA
miR-1	miR1GSP10	CATGATCAGCTGGGCCAAGATACATACTTC SEQ ID NO:47	miR-1RP	T+G+GAA+TG+TAAAGAAGT SEQ ID NO:48	Identical
miR-7	miR-7GSP10	CATGATCAGCTGGGCCAAGAAACAAAATC SEQ ID NO: 486	miR-7_RP6	T+GGAA+GACTTGTGATTTT SEQ ID NO: 487	one or more base pairs differ
miR-9*	mik-9*GSP	CATGATCAGCTGGGCCAAGAACTTTCGGTT SEQ ID NO:51	miR-9*RP	TAAA+GCT+AGATAACCG SEQ ID NO:52	Identical overlapping sequence, ends differ
miR-10a	miR-10aGSP	CATGATCAGCTGGGCCAAGACACAAATTCG SEQ ID NO:53	miR-10aRP	T+AC+CCTGTAGATCCG SEQ ID NO:54	Identical
miR-10b	miR-10b_GSP11	CATGATCAGCTGGGCCAAGAACACAAATTC G SEQ ID NO: 492	miR-10b_RP2	C+CC+TGT+AGAACCGAAT SEQ ID NO: 493	one or more base pairs differ
miR-15a	miR-15aGSP	CATGATCAGCTGGGCCAAGACCAATSEQ ID NO:57	miR-15aRP	T+AG+CAGCACATAATG SEQ ID NO:58	Identical
miR-15b	miR-15bGSP2	CATGATCAGCTGGGCCAAGATGTAAACCA SEQ ID NO:59	miR-15bRP	T+AG+CAGCACATCAT SEQ ID NO:60	Identical
miR-16	miR-16GSP2	CATGATCAGCTGGGCCAAGACGCCAATAT SEQ ID NO:61	miR-16RP	T+AG+CAGCACGTAAA SEQ ID NO:62	Identical
miR-17-3p	miR-17-3pGSP	CATGATCAGCTGGGCCAAGAACAAGTGCCC SEQ ID NO: 463	miR-17-3pRP	A+CT+GCAGTGAGGGC SEQ ID NO: 464	one or more base pairs differ
miR-17-5p	miR-17-5pGSP2	CATGATCAGCTGGGCCAAGAACTACCTGC SEQ ID NO: 65	miR-17-5pRP	C+AA+AGTGCTTACAGTG SEQ ID NO:66	Identical
miR-19a	miR-19aGSP2	CATGATCAGCTGGGCCAAGATCAGTTTTG SEQ ID NO:67	miR-19aRP	TG+TG+CAAATCTATGC SEQ ID NO:68	Identical
miR-19b	miR-19bGSP	CATGATCAGCTGGGCCAAGATCAGTTTTGC SEQ ID NO:69	miR-19bRP	TG+TG+CAAATCCATG SEQ ID NO:70	Identical

					Mouse microRNA as
Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	compared to Human microRNA
miR-20	miR-20GSP3	CATGATCAGCTGGGCCAAGACTACCTGC SEQ ID NO:71	miR-20RP	T+AA+AGTGCTTATAGTGCA SEQ ID NO:72	Identical
miR-21	miR-21GSP2	CATGATCAGCTGGGCCAAGATCAACATCA SEQ ID NO: 73	miR-21RP	T+AG+CTTATCAGACTGATG SEQ ID NO:74	Identical
miR-23a	miR-23aGSP	CATGATCAGCTGGGCCAAGAGGAAATCCCT SEQ ID NO:75	miR-23aRP	A+TC+ACATTGCCAGG SEQ ID NO:76	Identical
miR-23b	miR-23bGSP	CATGATCAGCTGGGCCAAGAGGTAATCCCT SEQ ID NO:77	miR-23bRP	A+TC+ACATTGCCAGG SEQ ID NO:78	Identical
mi.R-24	miR-24P5	CATGATCAGCTGGGCCAAGACTGTTCCTGC TG SEQ ID NO: 7	miR24-1,2R	TGG+CTCAGTTCAGC SEQ ID NO: 19	Identical
miR-25	miR-25GSP	CATGATCAGCTGGGCCAAGATCAGACCGAG SEQ ID NO:79	miR-25RP	C+AT+TGCACTTGTCTC SEQ ID NO:80	Identical
miR-26a	miR-26aGSP9	CATGATCAGCTGGGCCAAGAGCCTATCCT SEQ ID NO:81	miR-26aRP2	TT+CA+AGTAATCCAGGAT SEQ ID NO:82	Identical
miR-26b	miR-26bGSP9	CATGATCAGCTGGGCCAAGAAACCTATCC SEQ ID NO:83	miR-26bRP2	TT+CA+AGT+AATTCAGGAT SEQ ID NO:84	Identical
miR-27a	miR-27aGSP	CATGATCAGCTGGGCCAAGAGCGGAACTTA SEQ ID NO:85	mik-27akP	TT+CA+CAGTGGCTAA SEQ ID NO:86	Identical
miR-27b	miR-27bGSP	CATGATCAGCTGGGCCAAGAGCATASEQ ID NO:87	miR-27bRP	TT+CA+CAGTGGCTAA SEQ ID NO:88	Identical
miR-28	miR-28GSP	CATGATCAGCTGGGCCAAGACTCAATAGAC SEQ ID NO:89	miR-28RP	A+AG+GAGCTCACAGT SEQ ID NO:90	Identical
miR-29a	miR-29aGSP8	CATGATCAGCTGGGCCAAGAACCGATT SEQ ID NO:91	miR-29aRP2	T+AG+CACCATCTGAAAT SEQ ID NO:92	Identical
miR-29b	miR-29bGSP2	CATGATCAGCTGGGCCAAGAAACACTGAT SEQ ID NO:93	miR-29bRP2	T+AG+CACCATTTGAAATCAG SEQ ID NO:94	Identical

Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Mouse microRNA as compared to Human microRNA
miR-30a-5p	miR-30a-5pGSP	CATGATCAGCTGGGCCAAGACTTCCAGTCG	miR-30a-5pRP	T+GT+AAACATCCTCGAC	Identical
miR-30b	miR-30bGSP	CATGATCAGCTGGGCCAAGAAGCTGAGTGT	miR-30bRP	TGT+AAA+CATCCTACACT	Identical
		SEQ ID NO:97		SEQ ID NO:98	
miR-30c	miR-30cGSP	CATGATCAGCTGGGCCAAGAGCTGAGAGTG	miR-30cRP	TGT+AAA+CATCCTACACT	Identical
		SEQ ID NO:99		SEQ ID NO:100	
miR-30d	miR-30dGSP	CATGATCAGCTGGGCCAAGACTTCCAGTCG	miR-30dRP	T+GTAA+CATCCCCG	Identical
		SEQ ID NO:101		SEQ ID NO:102	
miR-30e-3p	miR-30e-3pGSP9	CATGATCAGCTGGGCCAAGAGCTGTAAAC	miR-30e-3pRP5	CTTT+CAGT+CGGATGTTT	Identical
		SEQ ID NO:103		SEQ ID NO:104	
miR-31	miR-31GSP	CATGATCAGCTGGGCCAAGACAGCTATGCC	miR-31RP	G+GC+AAGATGCTGGC	Identical overlapping sequence,
		SEQ ID NO:107		SEQ ID NO:108	ends differ
miR-32	miR-32GSP	CATGATCAGCTGGGCCAAGAGCAACTTAGT	miR-32RP	TATTG+CA+CATTACTAAG	Identical
į		SEQ ID NO:109		SEQ ID NO:110	
miR-33	miR-33GSP2	CATGATCAGCTGGGCCAAGACAATGCAAC	miR-33RP	G+TG+CATTGTAGTTGC	Identical
		SEQ ID NO:111		SEQ ID NO:112	
miR-34a	miR-34aGSP	CATGATCAGCTGGGCCAAGAAACAACCAGC	miR-34aRP	T+GG+CAGTGTCTTAG	Identical
		SEQ ID NO:113		SEQ ID NO:114	
miR-34b	miR-34bGSP	CATGATCAGCTGGGCCAAGACAATCAGCTA	miR-34bRP	TA+GG+CAGTGTAATT	one or more base pairs differ
		SEQ ID NO: 115		SEQ ID NO: 482	
miR-34c	miR-34cGSP	CATGATCAGCTGGGCCAAGAGCAATCAGCT	miR-34cRP	A+GG+CAGTGTAGTTA	Identical
		SEQ ID NO:117		SEQ ID NO:118	
miR-92	miR-92GSP	CATGATCAGCTGGGCCAAGACAGGCCGGGA	miR-92RP	T+AT+TGCACTTGTCCC	Identical
		SEQ ID NO:119		SEQ ID NO:120	
miR-93	miR-93GSP	CATGATCAGCTGGGCCAAGACTACCTGCAC	miR-93RP	AA+AG+TGCTGTTCGT	Identical overlapping sequence,
		SEQ ID NO:121		SEQ ID NO:122	ends differ
miR-96	miR-96GSP	CATGATCAGCTGGGCCAAGAGCAAAAATGT	miR-96RP	T+TT+GGCACTAGCAC	Identical overlapping sequence,
		SEQ ID NO:125		SEQ ID NO:126	ends differ

Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Mouse microRNA as compared to Human microRNA
miR-98	miR-98GSP	CATGATCAGCTGGGCCAAGAAACAATACAA SEQ ID NO:127	miR-98RP	TGA+GGT+AGTAAGTTG SEQ ID NO:128	Identical
miR-99a	miR-99aGSP	CATGATCAGCTGGGCCAAGACACAAGATCG SEQ ID NO:129	miR-99aRP	A+AC+CCGTAGATCCG SEQ ID NO:130	Identical overlapping sequence, ends differ
miR-99b	miR-99bGSP	CATGATCAGCTGGGCCAAGACGCAAGGTCG SEQ ID NO:131	miR-99bRP	C+AC+CCGTAGAACCG SEQ ID NO:132	Identical
miR-100	miR-100GSP	CATGATCAGCTGGGCCAAGACACAAGTTCG SEQ ID NO:133	miR-100RP	A+AC+CCGTAGATCCG SEQ ID NO:134	Identical
miR-101	miR-101GSP	CATGATCAGCTGGGCCAAGACTTCAGTTAT SEQ ID NO:135	miR-101RP	TA+CAG+TACTGTGATAACT SEQ ID NO:136	Identical
miR-103	miR-103GSP	CATGATCAGCTGGGCCAAGATCATAGCCCT SEQ ID NO:137	miR-103RP	A+GC+AGCATTGTACA SEQ ID NO:138	Identical
miR-106a	miR-106aGSP	CATGATCAGCTGGGCCAAGATACCTGCAC SEQ ID NO: 472	miR-106aRP	CAA+AG+TGCTAACAGTG SEQ ID NO: 473	one or more base pairs differ
miR-106b	miR-106bGSP	CATGATCAGCTGGGCCAAGAATCTGCACTG SEQ ID NO:143	miR-106bRP	T+AAAG+TGCTGACAGT SEQ ID NO:144	Identical
miR-107	miR-107GSP8	CATGATCAGCTGGGCCAAGATGATAGCC SEQ ID NO:145	miR-107RP2	A+GC+AGCATTGTACAG SEQ ID NO:146	Identical
miR-122a	miR-122aGSP	CATGATCAGCTGGGCCAAGAACAACACCA SEQ ID NO:147	miR-122aRP	T+GG+AGTGTGACAAT SEQ ID NO:148	Identical
miR-124a	miR-124aGSP	CATGATCAGCTGGGCCAAGATGGCATTCAC SEQ ID NO:149	miR-124aRP	T+TA+AGGCACGCGGT SEQ ID NO:150	Identical overlapping sequence, ends differ
miR-125a	miR-125aGSP	CATGATCAGCTGGGCCAAGACACAGGTTAA SEQ ID NO:151	miR-125aRP	T+CC+CTGAGACCCTT SEQ ID NO:152	Identical
miR-125b	miR-125bGSP	CATGATCAGCTGGGCCAAGATCACAAGTTA SEQ ID NO:153	miR-125bRP	T+CC+CTGAGACCCTA SEQ ID NO:154	Identical
miR-126	miR-126GSP	CATGATCAGCTGGGCCAAGAGCATTATTAC SEQ ID NO:155	miR-126RP	T+CG+TACCGTGAGTA SEQ ID NO:156	Identical

Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Mouse microRNA as compared to Human microRNA
miR-126*	miR-126*GSP3	CATGATCAGCTGGGCCAAGACGCGTACC SEQ ID NO:157	miR-126*RP	C+ATT+ATTA+CTTTTGGTACG SEQ ID NO:158	Identical
miR-127	miR-127GSP	CATGATCAGCTGGGCCAAGAAGCCAAGCTC SEQ ID NO:159	miR-127RP	T+CG+GATCCGTCTGA SEQ ID NO:160	Identical overlapping sequence, ends differ
miR-128a	miR-128aGSP	CATGATCAGCTGGGCCAAGAAAAAGAGACC SEQ ID NO:161	miR-128aRP	T+CA+CAGTGAACCGG SEQ ID NO:162	Identical
miR-128b	miR-128bGSP	CATGATCAGCTGGGCCAAGAGAAGAGACC SEQ ID NO:163	miR-128bRP	T+CA+CAGTGAACCGG SEQ ID NO:164	Identical
miR-130a	miR-130aGSP	CATGATCAGCTGGGCCAAGAATGCCCTTTT SEQ ID NO:167	mik-130aRP	C+AG+TGCAATGTTAAAAG SEQ ID NO:168	Identical.
miR-130b	miR-130bGSP	CATGATCAGCTGGGCCAAGAATGCCCTTTC SEQ ID NO:169	miR-130bRP	C+AG+TGCAATGATGA SEQ ID NO:170	Identical
miR-132	miR-132GSP	CATGATCAGCTGGGCCAAGACGACCATGGC SEQ ID NO:171	miR-132RP	T+AA+CAGTCTACAGCC SEQ ID NO:172	Identical
miR-133a	miR-133aGSP	CATGATCAGCTGGGCCAAGAACAGCTGGTT SEQ ID NO:173	miR-133aRP	T+TG+GTCCCCTTCAA SEQ ID NO:174	Identical
miR-133b	miR-133bGSP	CATGATCAGCTGGGCCAAGATAGCTGGTTG SEQ ID NO:175	mik-133bkP	T+TG+GTCCCTTCAA SEQ ID NO:176	Identical
miR-134	miR-134GSP	CATGATCAGCTGGGCCAAGACCCTCTGGTC SEQ ID NO:177	miR-134RP	T+GT+GACTGGTTGAC SEQ ID NO:178	Identical overlapping sequence, ends differ
miR-135a	miR-135aGSP	CATGATCAGCTGGGCCAAGATCACATAGGA SEQ ID NO:179	miR-135aRP	T+AT+GGCTTTTTATTCCT SEQ ID NO:180	Identical
miR-135b	miR-135bGSP	CATGATCAGCTGGGCCAAGACACATAGGAA SEQ ID NO:181	miR-135bRP	T+AT+GGCTTTTCATTCC SEQ ID NO:182	Identical
miR-136	miR-136GSP	CATGATCAGCTGGGCCAAGATCCATCATCA SEQ ID NO:183	miR-136RP	A+CT+CCATTTGTTTTGATG SEQ ID NO:184	Identical
miR-137	miR-137GSP	CATGATCAGCTGGGCCAAGACTACGCGTAT SEQ ID NO:185	miR-137RP	T+AT+TGCTTAAGAATACGC SEQ ID NO:186	Identical overlapping sequence, ends differ

Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Mouse microRNA as compared to Human microRNA
miR-138	miR-138GSP2	CATGATCAGCTGGGCCAAGACGGCCTGAT SEQ ID NO:187	miR-138RP	A+GC+TGGTGTTGTGA SEQ ID NO:188	Identical
miR-139	miR-139GSP	CATGATCAGCTGGGCCAAGAAGACACGTGC SEQ ID NO:189	miR-139RP	T+CT+ACAGTGCACGT SEQ ID NO:190	Identical
miR-140	miR-140GSP	CATGATCAGCTGGGCCAAGACTACCATAGG SEQ ID NO:191	miR-140RP	A+GT+GGTTTTACCCT SEQ ID NO:192	Identical overlapping sequence, ends differ
miR-141	miR-141GSP9	CATGATCAGCTGGGCCAAGACCATCTTA SEQ ID NO:193	miR-141RP2	TAA+CAC+TGTCTGGTAA SEQ ID NO:194	Identical
miR-142-3p	miR-142-3pGSP3	CATGATCAGCTGGGCCAAGATCCATAAA SEQ ID NO:195	miR-142-3pRP	TGT+AG+TGTTTCCTACT SEQ ID NO:196	Identical overlapping sequence, ends differ
miR-143	miR-143GSP8	CATGATCAGCTGGGCCAAGATGAGCTAC SEQ ID NO:197	miR-143RP2	T+GA+GATGAAGCACTG SEQ ID NO:198	Identical
miR-144	miR-144GSP2	CATGATCAGCTGGGCCAAGACTAGTACAT SEQ ID NO:199	miR-144RP	TA+CA+GTAT+AGATGATG SEQ ID NO:200	Identical
miR-145	miR-145GSP2	CATGATCAGCTGGGCCAAGAAAGGGATTC SEQ ID NO:201	miR-145RP	G+TC+CAGTTTTCCCA SEQ ID NO:202	Identical
miR-146	miR-146GSP3	CATGATCAGCTGGGCCAAGAACCCATG SEQ ID NO:203	miR-146RP	T+GA+GAACTGAATTCCA SEQ ID NO:204	Identical
miR-148a	miR-148aGSP2	CATGATCAGCTGGGCCAAGAACAAGTTC SEQ ID NO:207	miR-148aRP2	T+CA+GIGCACTACAGAACT SEQ ID NO:208	Identical
miR-148b	miR-148bGSP2	CATGATCAGCTGGGCCAAGAACAAAGTTC SEQ ID NO:209	miR-148bRP	T+CA+GTGCATCACAG SEQ ID NO:210	Identical
miR-149	miR-149GSP2	CATGATCAGCTGGGCCAAGAGGAGTGAAG SEQ ID NO:211	miR-149RP	T+CT+GGCTCCGTGTC SEQ ID NO:212	Identical
miR-150	miR-150GSP3	CATGATCAGCTGGGCCAAGACACTGGTA SEQ ID NO:213	miR-150RP	T+CT+CCCAACCCTTG SEQ ID NO:214	Identical
miR-151	miR-151GSP2	CATGATCAGCTGGGCCAAGACCTCAAGGA SEQ ID NO: 215	miR-151RP	A+CT+AGACTGAGGCTC SEQ ID NO: 477	one or more base pairs differ

					Mouse microRNA as
Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	compared to Human microRNA
miR-152	miR-152GSP2	CATGATCAGCTGGGCCAAGACCCAAGTTC	miR-152RP	T+CA+GTGCATGACAG SEO ID NO:218	Identical
miR-153	miR-153GSP2	CATGATCAGCTGGGCCAAGATCACTTTG	miR-153RP	TTG+CAT+AGTCACAAAA	Identical overlapping sequence, ends differ
miR-154	miR-154GSP9	CATGATCAGCTGGGCCAAGACGAAGGCAA	miR-154RP3	TA+GGTTA+TCCGTGTT SEQ ID NO:224	Identical
miR-155	miR-155GSP8	CATGATCAGCTGGGCCAAGACCCCTATC SEQ ID NO: 225	miR-155RP2	TT+AA+TGCTAATTGTGATAGG SEQ ID NO: 489	one or more base pairs differ
miR-181a	miR-181aGSP9	CATGATCAGCTGGGCCAAGAACTCACCGA SEQ ID NO:227	miR-181aRP2	AA+CATT+CAACGCTGTC SEQ ID NO:228	Identical
miR-181c	miR-181cGSP9	CATGATCAGCTGGGCCAAGAACTCACCGA SEQ ID NO:229	miR-181cRP2	AA+CATT+CAACCTGTCG SEQ ID NO:230	Identical
miR-182	miR-182*GSP	CATGATCAGCTGGGCCAAGATAGTTGGCAA SEQ ID NO:231	miR-182*RP	T+GG+TTCTAGACTTGC SEQ ID NO:232	Identical
miR-183	miR-183GSP2	CATGATCAGCTGGGCCAAGACAGTGAATT SEQ ID NO:235	miR-183RP	T+AT+GGCACTGGTAG SEQ ID NO:236	Identical
miR-184	miR-184GSP2	CATGATCAGCTGGGCCAAGAACCCTTATC SEQ ID NO:237	miR-184RP	T+GG+ACGGAGAACTG SEQ ID NO:238	Identical
miR-186	miR-186GSP9	CATGATCAGCTGGGCCAAGAAGCCCAAA SEQ ID NO:239	miR-186RP3	CA+AA+GAATT+CTCCTTTTGG SEQ ID NO:240	Identical
miR-187	miR-187GSP	CATGATCAGCTGGGCCAAGACGGCTGCAAC SEQ ID NO:241	miR-187RP	T+CG+TGTCTTGTGTT SEQ ID NO:242	Identical overlapping sequence, ends differ
miR-188	miR-188GSP	CATGATCAGCTGGGCCAAGAACCCTCCACC SEQ ID NO:243	miR-188RP	C+AT+CCCTTGCATGG SEQ ID NO:244	Identical
miR-189	miR-189GSP2	CATGATCAGCTGGGCCAAGAACTGATATC SEQ ID NO:245	miR-189RP	G+TG+CCTACTGAGCT SEQ ID NO:246	Identical
miR-190	miR-190GSP9	CATGATCAGCTGGGCCAAGAACCTAATAT SEQ ID NO:247	miR-190RP4	T+GA+TA+TGTTTGATATATTAG SEQ ID NO:248	Identical

Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Mouse microKNA as compared to Human microRNA
miR-191	miR-191GSP2	CATGATCAGCTGGGCCAAGAAGCTGCTTT SEO ID NO:249	miR-191RP2	C+AA+CGGAATCCCAAAAG SEQ ID NO:250	Identical
miR-192	miR-192GSP2	CATGATCAGCTGGGCCAAGAGGCTGTCAA SEQ ID NO:251	miR-192RP	C+TGA+CCTATGAATTGAC SEQ ID NO:252	Identical overlapping sequence, ends differ
miR-193	miR-193GSP9	CATGATCAGCTGGGCCAAGACTGGGACTT SEQ ID NO:253	miR-193RP2	AA+CT+GGCCTACAAAG SEQ ID NO:254	Identical
miR-194	mir194GSP8	CATGATCAGCTGGGCCAAGATCCACATG SEQ ID NO:255	mir194RP	TG+TAA+CAGCAACTCCA SEQ ID NO:256	Identical
miR-195	miR-195GSP9	CATGATCAGCTGGGCCAAGAGCCAATATT SEQ ID NO:257	miR-195RP3	T+AG+CAG+CACAGAAATA SEQ ID NO:258	Identical
miR-196a	miR-196aGSP	CATGATCAGCTGGGCCAAGACCAACAT SEQ ID NO:261	miR-196aRP	TA+GG+TAGTTTCATGTTG SEQ ID NO:262	Identical
miR-196b	miR-196bGSP	CATGATCAGCTGGGCCAAGACCAACAGG SEQ ID NO:259	miR-196bRP	TA+GGT+AGTTTCCTGT SEQ ID NO:260	Identical
miR-199a*	miR-199a*GSP2	CATGATCAGCTGGGCCAAGAAACCAATGT SEQ ID NO:267	miR-199a*RP	T+AC+AGTAGTCTGCAC SEQ ID NO:268	Identical
miR-199a	miR-199aGSP2	CATGATCAGCTGGGCCAAGAGAACAGGTA SEQ ID NO:269	miR-199aRP	C+CC+AGTGTTCAGAC SEQ ID NO:270	Identical
miR-199b	miR-199bGSP	CATGATCAGCTGGGCCAAGAGAACAGGTAG SEQ ID NO: 475	miR-199bRP	C+CC+AGTGTTTAGAC SEQ ID NO: 272	one or more base pairs differ
miR-200a	miR-200aGSP2	CATGATCAGCTGGGCCAAGAACATCGTTA SEQ ID NO:273	miR-200aRP	TAA+CAC+TGTCTGGT SEQ ID NO:274	Identical
miR-200b	miR-200bGSP2	CATGATCAGCTGGGCCAAGAGTCATCTT SEQ ID NO:275	miR-200bRP	TAATA+CTG+CCTGGTAAT SEQ ID NO:276	Identical
miR-203	miR-203GSP2	CATGATCAGCTGGGCCAAGACTAGTGGTC SEQ ID NO:279	miR-203RP	G+TG+AAATGTTTAGGACC SEQ ID NO:280	Identical overlapping sequence, ends differ
miR-204	miR-204GSP2	CATGATCAGCTGGGCCAAGAAGGCATAGG SEQ ID NO:281	miR-204RP	T+TC+CCTTTGTCATCC SEQ ID NO:282	Identical overlapping sequence, ends differ

1					Mouse microRNA as
Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	compared to Human microRNA
miR-205	miR-205GSP	CATGATCAGCTGGGCCAAGACAGACTCCGG	miR-205RP	T+CCTT+CATTCCACC	Identical
		SEQ 1D NO:283		SEQ ID NO: 204	
miR-206	mir206GSP7	CATGATCAGCTGGGCCAAGACCACACA	miR-206RP	T+G+GAA+TGTAAGGAAGTGT	Identical
		SEQ ID NO:285		SEQ ID NO:286	
miR-208	miR-208_GSP13	CATGATCAGCTGGGCCAAGAACAAGCTTTT	miR-208_RP4	ATAA+GA+CG+AGCAAAAAG	Identical
		rec SEQ ID NO:287		SEQ ID NO:288	
miR-210	miR-210GSP	CATGATCAGCTGGGCCAAGATCAGCCGCTG	miR-210RP	C+TG+TGCGTGTGACA	Identical
		SEQ ID NO:289		SEQ ID NO:290	
miR-211	miR-211GSP2	CATGATCAGCTGGGCCAAGAAGG	miR-211RP	T+TC+CCTTTGTCATCC	one or more base pairs differ
		SEQ ID NO: 491		SEQ ID NO: 292	
miR-212	miR-212GSP9	CATGATCAGCTGGGCCAAGAGGCCGTGAC	miR-212RP2	T+AA+CAGTCTCCAGTCA	Identical
		SEQ ID NO:293		SEQ ID NO:294	
miR-213	miR-213GSP	CATGATCAGCTGGGCCAAGAGGTACAATCA	miR-213RP	A+CC+ATCGACCGTTG	Identical
		SEQ ID NO:295		SEQ ID NO:296	
miR-214	miR-214GSP	CATGATCAGCTGGCCAAGACTGCCTGTCT	miR-214RP	A+CA+GCAGGCACAGA	Identical
		SEQ ID NO:297		SEQ ID NO:298	<u>s.</u>
miR-215	miR-215GSP2	CATGATCAGCTGGGCCAAGAGTCTGTCAA	miR-215RP	A+TGA+CCTATGATTTGAC	one or more base pairs differ
		SEQ ID NO: 299		SEQ ID NO: 469	
miR-216	miR-216GSP9	CATGATCAGCTGGGCCAAGACACAGTTGC	mir216RP	TAA+TCT+CAGCTGGCA	Identical
		SEQ ID NO:301		SEQ ID NO:302	
miR-217	miR-217GSP2	CATGATCAGCTGGGCCAAGAATCCAGTCA	miR-217RP2	T+AC+TGCATCAGGAACTGA	one or more base pairs differ
		SEQ ID NO:481		SEQ ID NO: 304	
miR-218	miR-218GSP2	CATGATCAGCTGGGCCAAGAACATGGTTA	miR-218RP	TTG+TGCTT+GATCTAAC	Identical
		SEQ ID NO:305		SEQ ID NO:306	
miR-221	miR-221GSP9	CATGATCAGCTGGGCCAAGAGAAACCCAG	miR-221RP	A+GC+TACATTGTCTGC	Identical overlapping sequence,
-		SEQ ID NO:309		SEQ ID NO:310	ends differ

Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Mouse microRNA as compared to Human microRNA
miR-222	miR-222GSP8	CATGATCAGCTGGGCCAAGAGAGCCCA SEQ ID NO:311	miR-222RP	A+GC+TACATCTGGCT SEQ ID NO:312	Identical
miR-223	miR-223GSP	CATGATCAGCTGGGCCAAGAGGGGTATTTG SEQ ID NO:313	miR-223RP	TG+TC+AGTTTGTCAAA SEQ ID NO:314	Identical
miR-224	miR-224GSP8	CATGATCAGCTGGGCCAAGATAAACGGA SEQ ID NO:315	miR-224RP2	C+AAG+TCACTAGTGGTT SEQ ID NO:316	Identical overlapping sequence, ends differ
miR-296	miR-296GSP9	CATGATCAGCTGGGCCAAGAACAGGATTG SEQ ID NO:317	miR-296RP2	A+GG+GCCCCCTCAA SEQ ID NO:318	Identical
miR-299	miR-299GSP9	CATGATCAGCTGGGCCAAGAATGTATGTG SEQ ID NO:319	miR-299RP	T+GG+TTTACCGTCCC SEQ ID NO:320	Identical
miR-301	miR-301GSP	CATGATCAGCTGGGCCAAGAGCTTTGACAA SEQ ID NO:321	miR-301RP	C+AG+TGCAATAGTATTGT SEQ ID NO:322	
miR-302a	miR-302aGSP	CATGATCAGCTGGGCCAAGATCACCAAAAC SEQ ID NO:325	miR-302aRP	T+AAG+TGCTTCCATGT SEQ ID NO:326	Identical
miR-320	miR-320_GSP8	CATGATCAGCTGGGCCAAGATTCGCCCT SEQ ID NO:337	miR-320_RP3	AAAA+GCT+GGGTTGAGAGG SEQ ID NO:338	Identical
miR-323	miR-323GSP	CATGATCAGCTGGGCCAAGAAGAGGTCGAC SEQ ID NO:339	miR-323RP	G+CA+CATTACACGGT SEQ ID NO:340	Identical
miR-324-3p	miR-324-3pGSP	CATGATCAGCTGGGCCAAGACCAGCACCAC SEQ ID NO:341	miR-324-3pRP	C+CA+CTGCCCCAGGT SEQ ID NO:342	Identical
miR-324-5p	miR-324-5pGSP	CATGATCAGCTGGGCCAAGAACACCAATGC SEQ ID NO:343	miR-324-5pRP	C+GC+ATCCCTAGGG SEQ ID NO:344	Identical overlapping sequence, ends differ
miR-325	miR-325GSP	CATGATCAGCTGGGCCAAGAACACTTACTG	miR-325RP	C+CT+AGTAGGTGCTC SEQ ID NO: 476	one or more base pairs differ
miR-326	miR-326GSP	CATGATCAGCTGGGCCAAGACTGGAGGAAG	miR-326RP	C+CT+CTGGGCCCTTC SEQ ID NO:348	Identical overlapping sequence, ends differ
miR-328	miR-328GSP	CATGATCAGCTGGGCCAAGAACGGAAGGGC SEQ ID NO:349	miR-328RP	C+TG+GCCCTCTCTGC SEQ ID NO:350	Identical

microRNA:			9		Mouse microRNA as
	Extension Primer Name	Extension Primer Sequence	Primer Name	Reverse Primer Sequence	microRNA
	miR-330GSP	TGGGCCAAGATCTCTGCAGG	miR-330RP	G+CA+AAGCACAGGGC	one or more base pairs differ
		SEQ ID NO: 351		SEQ ID NO: 4/8	
miR-331 mi	miR-331GSP	CATGATCAGCTGGGCCAAGATTCTAGGATA	miR-331RP	G+CC+CCTGGGCCTAT	Identical
		SEQ ID NO:353		SEQ ID NO:354	
miR-337 mj	miR-337GSP	CATGATCAGCTGGGCCAAGAAAAGGCATCA	miR-337RP	T+TC+AGCTCCTATATG	one or more base pairs differ
		SEQ ID NO:355		SEQ ID NO: 490	
miR-338 mi	miR-338GSP	CATGATCAGCTGGGCCAAGATCAACAAAT	miR-338RP2	T+CC+AGCATCAGTGATTT	Identical
		SEQ ID NO:357		SEQ ID NO:358	
miR-339 mi	miR-339GSP9	CATGATCAGCTGGGCCAAGATGAGCTCCT	miR-339RP2	T+CC+CTGTCCTCCAGG	Identical
		SEQ ID NO:359		SEQ ID NO:360	
miR-340 mi	miR-340GSP	CATGATCAGCTGGGCCAAGAGGCTATAAAG	miR-340RP	TC+CG+TCTCAGTTAC	Identical
		SEQ ID NO:361		SEQ ID NO:362	
miR-342 mj	miR-342GSP3	CATGATCAGCTGGGCCAAGAGACGGGTG	miR-342RP	T+CT+CACACAGAAATCG	Identical
		SEQ ID NO:363		SEQ ID NO:364	•
miR-345 mi	miR-345GSP	CATGATCAGCTGGGCCAAGAGCACTGGACT	miR-345RP	T+GC+TGACCCCTAGT	one or more base pairs differ
		SEQ ID NO: 484		SEQ ID NO: 485	
miR-346 mj	miR-346GSP	CATGATCAGCTGGGCCAAGAAGAGGCAGGC	miR-346RP	T+GT+CTGCCCGAGTG	one or more base pairs differ
		SEQ ID NO: 367		SEQ ID NO: 488	
miR-363 m	miR-363 GSP10	CATGATCAGCTGGGCCAAGATACAGATGGA	miR-363RP	AAT+TG+CAC+GGTATCC	Identical
		SEQ ID NO:369		SEQ ID NO:370	
miR-370 mi	miR-370GSP	CATGATCAGCTGGGCCAAGACCAGGTTCCA	miR-370RP	G+CC+TGCTGGGGTGG	Identical overlapping sequence,
		SEQ ID NO:375		SEQ ID NO:376	ends differ
miR-375 mi	miR-375GSP	CATGATCAGCTGGGCCAAGATCACGCGAGC	miR-375RP	TT+TG+TTCGTTCGGC	Identical
		SEQ ID NO:387		SEQ ID NO:388	
miR-376a mi	miR-376aGSP3	CATGATCAGCTGGGCCAAGAACGTGGAT	miR-376aRP2	A+TCGTAGA+GGAAAATCCAC	one or more base pairs differ
		SEQ ID NO: 467		SEQ ID NO: 468	
miR-378 m	miR-378GSP	CATGATCAGCTGGGCCAAGAACACAGGACC	miR-378RP	C+TC+CTGACTCCAGG	Identical
		SEQ ID NO:391		SEQ ID NO:392	

Monse Target	Extension		Reverse		Mouse microRNA as compared to Human
microRNA:	Primer Name	Extension Primer Sequence	Primer Name	Reverse Primer Sequence	microRNA
miR-379	miR-379_GSP7	CATGATCAGCTGGGCCAAGATACGTTC SEQ ID NO:393	miR-379RP2	T+GGT+AGACTATGGAACG SEQ ID NO:394	Identical overlapping sequence, ends differ
miR-380-5p	miR-380-5pGSP	CATGATCAGCTGGGCCAAGAGCGCATGTTC SEQ ID NO:395	miR-380-5pRP	T+GGT+TGACCATAGA SEQ ID NO:396	Identical
miR-380-3p	miR-380-3pGSP	CATGATCAGCTGGGCCAAGAAAGATGTGGA SEQ ID NO: 395	miR-380-3pRP	TA+TG+TAGTATGGTCCACA SEQ ID NO: 483	one or more base pairs differ
miR-381	miR-381GSP2	CATGATCAGCTGGGCCAAGAACAGAGGC SEQ ID NO:399	miR-381RP2	TATA+CAA+GGGCAAGCT SEQ ID NO:400	Identical
miR-382	miR-382GSP	CATGATCAGCTGGGCCAAGACGAATCCACC SEQ ID NO:401	miR-382RP	G+AA+GTTGTTCGTGGT SEQ ID NO:402	Identical
miR-383	miR-383GSP	CATGATCAGCTGGGCCAAGAAGCCACAGTC SEQ ID NO:465	miR-383RP2	A+GATC+AGAAGGTGACTGT SEQ ID NO: 466	one or more base pairs differ
miR-384	miR-384_GSP9	CATGATCAGCTGGGCCAAGATGTGAACAA SEQ ID NO:470	miR-384_RP5	ATT+CCT+AG+AAATTGTTC SEQ ID NO: 471	one or more base pairs differ
miR-410	miR-410 GSP9	CATGATCAGCTGGGCCAAGAACAGGCCAT SEQ ID NO:405	miR-410RP	AA+TA+TAA+CA+CAGATGGC SEQ ID NO:406	Identical
miR-412	miR-412 GSP10	CATGATCAGCTGGGCCAAGAACGCCTAGTG SEQ ID NO:407	miR-412RP	A+CTT+CACCTGGTCCACTA SEQ ID NO:408	Identical
miR-424	miR-424GSP	CATGATCAGCTGGGCCAAGATCCAAAACAT SEQ ID NO:474	miR-424RP2	C+AG+CAGCAATTCATGTTTT SEQ ID NO: 414	one or more base pairs differ
miR-425	miR-425GSP	CATGATCAGCTGGGCCAAGAGGCGGACACG SEQ ID NO:417	miR-425RP	A+TC+GGGAATGTCGT SEQ ID NO:418	Identical
miR-429	miR-429_GSP11	CATGATCAGCTGGGCCAAGAACGCCATTACC SEQ ID NO: 479	miR-429RP5	T+AATAC+TG+TCTGGTAATG SEQ ID NO: 480	one or more base pairs differ
miR-431	miR-431 GSP10	CATGATCAGCTGGGCCAAGATGCATGACGG SEQ ID NO:421	miR-431RP	T+GT+CTTGCAGGCCG SEQ ID NO:422	Identical overlapping sequence, ends differ
miR-448	miR-448GSP	CATGATCAGCTGGGCCAAGAATGGGACATC SEQ ID NO:423	miR-448RP	TTG+CAIA+TGIAGGAIG SEQ ID NO:424	Identical

Mouse Target microRNA:	Extension Primer Name	Extension Primer Sequence	Reverse Primer Name	Reverse Primer Sequence	Mouse microRNA as compared to Human microRNA
miR-449	miR-449GSP10	CATGATCAGCTGGGCCAAGAACCAGCTAAC SEQ ID NO:425	miR-449RP2	T+GG+CAGTGTATTGTTAGC SEQ ID NO:426	Identical
miR-450	miR-450GSP	CATGATCAGCTGGGCCAAGATATTAGGAAC SEQ ID NO:427	miR-450RP	TTTT+TG+CGATGTGTT SEQ ID NO:428	Identical
miR-451	miR-451 GSP10	CATGATCAGCTGGGCCAAGAAACTCAGTA SEQ ID NO:429	miR-451RP	AAA+CCG+TTA+CCATTACTGA SEQ ID NO:430	Identical overlapping sequence, ends differ
let7a	let7a-GSP2	CATGATCAGCTGGGCCAAGAAACTATAC SEQ ID NO:431	let7a-RP	T+GA+GGTAGTAGGTTG SEQ ID NO:432	Identical overlapping sequence, ends differ
let7b	let7b-GSP2	CATGATCAGCTGGGCCAAGAAACCACAC SEQ ID NO: 433	let7b-RP	T+GA+GGTAGTTG SEQ ID NO:432	Identical
let7c	let7c-GSP2	CATGATCAGCTGGGCCAAGAACCATAC SEQ ID NO:434	let7c-RP	T+GA+GGTAGTTG SEQ ID NO:432	Identical
let7d	let7d-GSP2	CATGATCAGCTGGGCCAAGAACTATGCA SEQ ID NO: 435	let7d-RP	A+GA+GGTAGGTTG SEQ ID NO:436	Identical
let7e	let7e-GSP2	CATGATCAGCTGGGCCAAGAACTATACA SEQ ID NO:437	let7e-RP	T+GA+GGTAGGAGGTTG SEQ ID NO:438	Identical
let7f	let7f-GSP2	CATGATCAGCTGGGCCAAGAAACTATAC SEQ ID NO:439	let7f-RP	T+GA+GGTAGATTG SEQ ID NO:440	Identical overlapping sequence, ends differ
let7g	let7g-GSP2	CATGATCAGCTGGGCCAAGAACTGTACA SEQ ID NO:441	let7g-RP	T+GA+GGTAGTTTG SEQ ID NO:442	Identical
let7i	let7i-GSP2	CATGATCAGCTGGGCCAAGAACAGCACA	let7i-RP	T+GA+GGTAGTTTG SEQ ID NO:444	Identical

EXAMPLE 5

This Example describes the detection and analysis of expression profiles for three microRNAs in total RNA isolated from twelve different tissues using methods in accordance with an embodiment of the present invention.

Methods: Quantitative analysis of miR-1, miR-124 and miR-150 microRNA templates was determined using 0.5 μg of First Choice total RNA (Ambion, Inc.) per 10 μl primer extension reaction isolated from the following tissues: brain, heart, intestine, kidney, liver, lung, lymph, ovary, skeletal muscle, spleen, thymus and uterus. The primer extension enzyme and quantitative PCR reactions were carried out as described above in EXAMPLE 3, using the following PCR primers:

miR-1 template:

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extension primer: CATGATCAGCTGGGCCAAGATACATACTTC (SEQ ID NO: 47)

reverse primer: T+G+GAA+TG+TAAAGAAGT (SEQ ID NO: 48)

15 forward primer: CATGATCAGCTGGGCCAAGA (SEQ ID NO: 13)

miR-124 template:

extension primer: CATGATCAGCTGGGCCAAGATGGCATTCAC (SEQ ID NO: 149)

reverse primer: T+TA+AGGCACGCGGT (SEQ ID NO: 150)

20 forward primer: CATGATCAGCTGGGCCAAGA (SEQ ID NO: 13)

miR-150 template:

extension primer: CATGATCAGCTGGGCCAAGACACTGGTA (SEQ ID NO: 213)

reverse primer: T+CT+CCCAACCCTTG (SEQ ID NO: 214)

25 forward primer: CATGATCAGCTGGGCCAAGA (SEQ ID NO: 13)

Results: The expression profiles for miR-1, miR-124 and miR-150 are shown in FIGURE 3A, 3B, and 3C, respectively. The data in FIGURES 3A-3C are presented in units of microRNA copies per 10 pg of total RNA (y-axis). These units were chosen since human cell lines typically yield \leq 10 pg of total RNA per cell. Hence the data shown are estimates of microRNA copies per cell. The numbers on the x-axis correspond

to the following tissues: (1) brain, (2) heart, (3) intestine, (4) kidney, (5) liver, (6) lung, (7) lymph, (8) ovary, (9) skeletal muscle, (10) spleen, (11) thymus and (12) uterus.

Consistent with previous reports, very high levels of striated muscle-specific expression were found for miR-1 (as shown in FIGURE 3A), and high levels of brain expression were found for miR-124 (as shown in FIGURE 3B) (see Lagos-Quintana et al., RNA 9:175-179, 2003). Quantitative analysis reveals that these microRNAs are present at tens to hundreds of thousands of copies per cell. These data are in agreement with quantitative Northern blot estimates of miR-1 and miR-124 levels (see Lim et al., Nature 433:769-773, 2005). As shown in FIGURE 3C, miR-150 was found to be highly expressed in the immune-related lymph node, thymus and spleen samples which is also consistent with previous findings (see Baskerville et al., RNA 11:241-247, 2005).

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While the preferred embodiment of the invention has been illustrated and described, it will be appreciated that various changes can be made therein without departing from the spirit and scope of the invention.